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(54) Title: PLANTS HAVING MODIFIED GROWTH CHARACTERISTICS AND A METHOD FOR MAKING THE SAME

(57) Abstract: The present invention concerns a method for modifying the growth characteristics of plants by modifying expression in a plant of a nucleic acid sequence encoding a 2xC2H2 zinc finger protein and/or modifying level and/or activity in a plant of a 2xC2H2 zinc finger protein. The invention also relates to transgenic plants having modified growth characteristics, which plants have modified expression of a nucleic acid encoding a 2xC2H2 zinc finger protein. For example yield of crop plants are improved by the methods of the present invention.

Plants having modified growth characteristics and a method for making the same

The present invention concerns a method for modifying plant growth characteristics. More specifically, the present invention concerns a method for modifying the growth characteristics of a plant by modifying expression of a nucleic acid encoding a zinc finger protein and/or by modifying the level and/or activity of a zinc finger protein in a plant, which zinc finger protein has two zinc finger domains of the type C2H2 (2xC2H2). The present invention also concerns plants having modified expression of a nucleic acid encoding a 2xC2H2 zinc finger protein and/or modified levels and/or activity of a 2xC2H2 zinc finger protein, which plants have modified growth characteristics relative to corresponding wild type plants.

Given the ever-increasing world population, it remains a major goal of agricultural research to improve the efficiency of agriculture. Conventional means for crop and horticultural improvements utilise selective breeding techniques to identify plants having desirable characteristics. However, such selective breeding techniques have several drawbacks, namely that these techniques are typically labour intensive and result in plants that often contain heterogeneous genetic components that may not always result in the desirable trait being passed on from parent plants. Advances in molecular biology have allowed mankind to modify the germplasm of animals and plants in a specific and controlled way. Genetic engineering of plants entails the isolation and manipulation of genetic material (typically in the form of DNA or RNA) and the subsequent introduction of that genetic material into a plant. Such technology has led to the development of plants having various improved economic, agronomic or horticultural traits. A trait or growth characteristic of particular economic interest is high yield. Yield is normally defined as the measurable produce of economic value from a crop. This may be defined in terms of quantity and/or quality. Other important growth characteristics include modified architecture, modified growth rate, among others.

The ability to influence one or more of the abovementioned growth characteristics, would have many applications in areas such as crop enhancement, plant breeding, production of ornamental plants, arboriculture, horticulture, forestry, production of algae or plants (for example for use as bioreactors, for the production of substances such as pharmaceuticals, antibodies, or vaccines, or for the bioconversion of organic waste or for use as fuel in the case of high-yielding algae and plants).

The term "zinc finger" describes a nucleic acid-binding domain in a protein that is folded around a tetrahedrally coordinated Zinc ion (Miller et al. 1985. EMBO, 4, 1609-1614). The amino acids that coordinate the zinc ion, are always cystein or histidine residues, however, diversity occurs in the sequence and length of the zinc finger domain. Zinc finger proteins may

5 contain several zinc finger domains of the same or different type. Further variability is encountered in nature by association of zinc finger domains with other domains. For example, some zinc finger proteins are found in association with ring finger or coil-coil domains, to form a so-called tripartite domain. There are several types of zinc fingers, such as C2H2, C2HC, C2C2. C2H2 is known as the classical zinc finger domain. There are typically two criteria used

10 to classify zinc finger proteins, the first being the type of zinc finger and the second being the number of zinc fingers present in the protein. Zinc finger proteins having a single C2H2 domain have been characterised, for example Superman from *Arabidopsis* and Ramosa I from maize. A well-characterised zinc finger protein having three C2H2 domains is the Indeterminate 1 protein from Maize. Although the first report of this gene (Colasanti et al., Cell. 1998 May 15;93(4):593-603) only mentions the presence of two zinc finger domains, a more sophisticated analysis, using pFAM domain search, revealed the presence of three C2H2 zinc finger domains. Also known are zinc-finger proteins having only two C2H2 domains, for example ZAT10 (STZ) and SCOF-1. This subset of plant zinc finger proteins having two C2H2 domains have been implicated in plant responses to various stresses (Sakamoto et al., Gene 20 248 (1-2) 23-32 (2000)). Both STZ and SCOF-1 have been used to enhance abiotic stress tolerance. When over-expressed, STZ has been reported to increase salt tolerance in yeast (Lippuner et al., J Biol Chem. 271 (22) 12859-12866 (1996)) and over-expression of the SCOF-1 gene under control of the CaMV 35 S promoter has been reported to enhance cold tolerance in *Arabidopsis thaliana* (Kim et al., Plant J. 25 (3) 247-259 (2001)). Reports of plants

25 having modified expression of a zinc finger encoding gene (whether the zinc finger gene is mutated, over-expressed or otherwise) describe plants having abnormal growth characteristics, none of which (with the exception of cold stress tolerance in transgenic plants expressing SCOF-1) are desirable for crops or describe effects that are only detectable under particular stress conditions.

30 It has now been found that modifying expression in a plant of a 2xC2H2 zinc finger gene and/or modifying the level and/or activity in a plant of a 2xC2H2 zinc finger protein gives plants having modified growth characteristics. In particular it has been found that introduction into a plant of a 2xC2H2 zinc finger nucleic acid gives plants modified growth characteristics, such as

35 increased yield, modified leaf architecture and altered cycle time, each relative to wild type plants.

Therefore according to one embodiment of the present invention there is provided a method for modifying the growth characteristics of a plant, comprising modifying expression in a plant of a nucleic acid encoding a 2xC2H2 zinc finger protein and/or modifying level and/or activity in a plant of a 2xC2H2 zinc finger protein.

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The term "modifying" as used herein is taken to mean enhancing, decreasing and/or changing in place and/or time. Modifying expression of a nucleic acid encoding a 2xC2H2 zinc finger protein or modifying the level and/or activity of the 2xC2H2 zinc finger protein itself encompasses altered expression of a gene and/or altered level and/or activity of a gene

10 product, namely a polypeptide, in specific cells or tissues, when compared to expression, level and/or activity of a 2xC2H2 zinc finger gene or protein in corresponding wild-type plants. The modified gene expression may result from modified expression of an endogenous 2xC2H2 zinc finger gene and/or may result from modified expression of a 2xC2H2 zinc finger gene previously introduced into a plant. Similarly, modified levels and/or activity of a 2xC2H2 zinc finger protein may be due to modified expression of an endogenous 2xC2H2 zinc finger nucleic acid/gene and/or due to modified expression of a 2xC2H2 zinc finger nucleic acid/gene previously introduced into a plant. Modified expression of a gene/nucleic acid and/or modified level and/or activity of a gene product/protein may be effected, for example, by chemical means and/or recombinant means.

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Therefore there is provided by the present invention, a method for modifying the growth characteristics of a plant, comprising modifying expression, level and/or activity of a 2xC2H2 zinc finger gene or protein by recombinant means and/or by chemical means.

25 Advantageously, modifying expression of a nucleic acid encoding a 2xC2H2 zinc finger protein and/or modifying level and/or activity of the 2xC2H2 zinc finger protein itself may be effected by chemical means, i.e. by exogenous application of one or more compounds or elements capable of modifying activity of the 2xC2H2 zinc finger protein and/or capable of modifying expression of a 2xC2H2 zinc finger gene (which may be either an endogenous gene or a
30 transgene introduced into a plant). The term "exogenous application" as defined herein is taken to mean the contacting or administering of a suitable compound or element to a plant. The compound or element may be exogenously applied to a plant in a form suitable for plant uptake (such as through application to the soil for uptake via the roots, or in the case of some plants by applying directly to the leaves, for example by spraying). The exogenous application
35 may take place on wild-type plants or on transgenic plants that have previously been transformed with a 2xC2H2 zinc finger nucleic acid/gene or other transgene.

Suitable compounds or elements for exogenous application include 2xC2H2 zinc finger proteins or 2xC2H2 zinc finger nucleic acids. Alternatively, exogenous application of compounds or elements capable of modifying levels of factors that directly or indirectly activate or inactivate a 2xC2H2 zinc finger protein will also be suitable in practising the invention. Also 5 included are antibodies that can recognise or mimic the function of 2xC2H2 zinc finger proteins. Such antibodies may comprise "plantibodies", single chain antibodies, IgG antibodies and heavy chain camel antibodies, as well as fragments thereof.

Additionally or alternatively, the resultant effect may also be achieved by the exogenous 10 application of an interacting protein or activator or an inhibitor of a 2xC2H2 zinc finger gene/gene product. Additionally or alternatively, the compound or element may be a mutagenic substance, such as a chemical selected from any one or more of: N-nitroso-N-ethylurea, ethylene imine, ethyl methanesulphonate and diethyl sulphate. Mutagenesis may also be achieved by exposure to ionising radiation, such as X-rays or gamma-rays or ultraviolet light. 15 Methods for introducing mutations and for testing the effect of mutations (such as by monitoring gene expression and/or protein activity) are well known in the art.

Additionally or alternatively, and according to a preferred embodiment of the present invention, modifying expression of a nucleic acid encoding a 2xC2H2 zinc finger protein and/or modifying 20 level and/or activity of the 2xC2H2 zinc finger protein may be effected by recombinant means. Such recombinant means may comprise a direct and/or indirect approach for modifying expression of a nucleic acid and/or level and/or activity of a protein.

For example, an indirect approach may comprise introducing, into a plant, a nucleic acid 25 capable of modifying expression of the gene in question (a gene encoding a 2xC2H2 zinc finger protein) and/or capable of modifying the level and/or activity of the protein in question (a 2xC2H2 zinc finger protein). Examples of such nucleic acids to be introduced into a plant include nucleic acids encoding transcription factors or activators or inhibitors that bind to the promoter of a 2xC2H2 zinc finger gene or that interact with a 2xC2H2 zinc finger protein. 30 Methods to test these types of interactions and methods for isolating nucleic acids encoding such interactors include yeast one-hybrid or yeast two-hybrid screens in which the 2xC2H2 zinc finger gene/protein is used as bait. One example of such a transcription regulator is LOS2, described as a transcription regulator for the STZ gene. Therefore, the method of the invention may also be performed using LOS2, wherein expression of a 2xC2H2 zinc finger gene may be 35 increased or further increased by decreasing expression of LOS2 in plants.

Also encompassed by an indirect approach for modifying expression of a 2xC2H2 zinc finger gene and/or for modifying level and/or activity of a 2xC2H2 zinc finger protein is the provision of, or the inhibition or stimulation of regulatory sequences that drive expression of a native 2xC2H2 zinc finger gene or transgene. Such regulatory sequences may be introduced into a 5 plant. For example, the regulatory sequence to be introduced into a plant may be a promoter capable of driving expression of an endogenous 2xC2H2 zinc finger gene.

A further indirect approach for modifying expression of a 2xC2H2 zinc finger gene and/or for modifying level and/or activity of a 2xC2H2 zinc finger protein in a plant encompasses 10 modifying levels in a plant of a factor capable of interacting with a zinc finger protein. Such factors may include ligands of a 2xC2H2 zinc finger protein. Therefore, the present invention also provides a method for modifying growth characteristics of a plant, comprising modifying expression of a gene coding for a protein which is a natural ligand of a 2xC2H2 zinc finger protein. Furthermore, the present invention also provides a method for modifying growth 15 characteristics of a plant, comprising modifying expression of a gene coding for a protein which is a natural target/substrate of a 2xC2H2 zinc finger protein. Examples of such targets/substrates include stretches of DNA that are bound by the zinc-finger domains.

A direct and preferred approach on the other hand comprises introducing into a plant a nucleic 20 acid encoding a 2xC2H2 zinc finger protein or a portion thereof or sequences capable of hybridising therewith, which nucleic acid preferably encodes a 2xC2H2 zinc finger protein or a homologue, derivative or active fragment thereof. The nucleic acid may be introduced into a plant by, for example, transformation.

25 Therefore, there is provided a method for modifying growth characteristics of a plant, comprising introducing into a plant a 2xC2H2 zinc finger nucleic acid or a portion thereof.

The 2xC2H2 zinc finger nucleic acid may be derived (either directly or indirectly (if 30 subsequently modified)) from any source provided that the sequence, when expressed in a plant, leads to modified expression of a 2xC2H2 zinc finger-encoding nucleic acid/gene and/or modified level and/or activity of a 2xC2H2 zinc finger protein. The 2xC2H2 zinc finger gene or protein may be wild type, i.e. the native or endogenous nucleic acid or polypeptide. Alternatively, it may be a protein or nucleic acid derived from the same or another species. The 35 nucleic acid/gene may then be introduced into a plant as a transgene, for example by transformation.

The nucleic acid may be isolated from a bacteria, yeast or fungi, or from a plant, algae, insect or animal (including human) source. This nucleic acid may be substantially modified from its native form in composition and/or genomic environment through deliberate human manipulation. The nucleic acid is preferably obtained from a plant, whether from the same 5 plant species in which it is to be introduced or whether from a different plant species. Further preferably, the nucleic acid is from a dicot, preferably from the family *Brassicaceae*, further preferably from *Arabidopsis thaliana*. More preferably, the nucleic acid is essentially similar to a nucleic acid as represented by SEQ ID NO 1, or a portion of SEQ ID NO 1, or a nucleic acid capable of hybridising therewith or is a nucleic acid encoding an amino acid sequence 10 essentially similar to an amino acid as represented by SEQ ID NO 2, or a homologue, derivative or active fragment thereof.

Advantageously, the methods according to the invention may also be practised using variant 2xC2H2 zinc finger nucleic acids and variant 2xC2H2 zinc finger amino acids, preferably 15 wherein the variant nucleic acids are variants of SEQ ID NO 1 and wherein the variant amino acids are variants of SEQ ID NO 2. Examples of variant sequences suitable in performing the methods of the invention include:

- (i) Functional portions of a 2xC2H2 zinc finger nucleic acid/gene;
- (ii) Sequences capable of hybridising with a 2xC2H2 zinc finger nucleic acid/gene;
- 20 (iii) Alternative splice variants of a 2xC2H2 zinc finger nucleic acid/gene;
- (iv) Allelic variants of a 2xC2H2 zinc finger nucleic acid/gene;
- (v) Homologues, derivatives and active fragments of a 2xC2H2 zinc finger protein.

The abovementioned variants may also be described as being "essentially similar" to a 25 2xC2H2 zinc finger nucleic acid/gene, particularly to the 2xC2H2 zinc finger encoding nucleic acid of SEQ ID NO 1, or essentially similar to a 2xC2H2 zinc finger amino acid/protein, particularly that of SEQ ID NO 2. The term "essentially similar to" also includes variants of SEQ ID NO 1 in the form of a complement, DNA, RNA, cDNA or genomic DNA. The variant nucleic acid encoding a 2xC2H2 zinc finger protein or the variant of a 2xC2H2 zinc finger 30 protein may be synthesized in whole or in part, it may be a double-stranded nucleic acid or a single-stranded nucleic acid. Also, the term encompasses a variant due to the degeneracy of the genetic code; a family member of the gene or protein; and variants that are interrupted by one or more intervening sequences.

35 An example of a variant 2xC2H2 zinc finger nucleic acid is a functional portion of a 2xC2H2 zinc-finger gene. Advantageously, the method according to the present invention may also be practised using portions of a DNA or nucleic acid encoding a 2xC2H2 zinc finger protein. A

functional portion refers to a piece of DNA derived or prepared from an original (larger) DNA molecule, which DNA portion, when expressed in a plant, gives plants having modified growth characteristics. The portion may comprise many genes, with or without additional control elements or may contain spacer sequences. The portion may be made by making one or 5 more deletions and/or truncations to the nucleic acid. Techniques for introducing truncations and deletions into a nucleic acid are well known in the art. Portions suitable for use in the methods according to the invention may readily be determined by following the methods described in the Examples section by simply substituting the sequence used in the actual Example with the portion to be tested for functionality.

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An example of a further variant 2xC2H2 zinc finger nucleic acid is a sequence that is capable of hybridising to a 2xC2H2 zinc finger nucleic acid, for example to any of SEQ ID NO 1, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 41, 43, 45, 47 or 49. Advantageously, the methods according to the present invention may also be practised using these variants.

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Hybridising sequences suitable for use in the methods according to the invention may readily be determined for example by following the methods described in the Examples section by simply substituting the sequence used in the actual Example with the hybridising sequence.

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The term "hybridisation" as defined herein is a process wherein substantially homologous complementary nucleotide sequences anneal to each other. The hybridisation process can occur entirely in solution, i.e. both complementary nucleic acids are in solution. Tools in molecular biology relying on such a process include the polymerase chain reaction (PCR; and all methods based thereon), subtractive hybridisation, random primer extension, nuclease S1 mapping, primer extension, reverse transcription, cDNA synthesis, differential display of RNAs,

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and DNA sequence determination. The hybridisation process can also occur with one of the complementary nucleic acids immobilised to a matrix such as magnetic beads, Sepharose beads or any other resin. Tools in molecular biology relying on such a process include the isolation of poly (A+) mRNA. The hybridisation process can furthermore occur with one of the complementary nucleic acids immobilised to a solid support such as a nitro-cellulose or nylon

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membrane or immobilised by e.g. photolithography to, for example, a siliceous glass support (the latter known as nucleic acid arrays or microarrays or as nucleic acid chips). Tools in molecular biology relying on such a process include RNA and DNA gel blot analysis, colony hybridisation, plaque hybridisation, *in situ* hybridisation and microarray hybridisation. In order to allow hybridisation to occur, the nucleic acid molecules are generally thermally or chemically

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denatured to melt a double strand into two single strands and/or to remove hairpins or other secondary structures from single stranded nucleic acids. The stringency of hybridisation is influenced by conditions such as temperature, salt concentration and hybridisation buffer

composition. High stringency conditions for hybridisation include high temperature and/or low salt concentration (salts include NaCl and Na₃-citrate) and/or the inclusion of formamide in the hybridisation buffer and/or lowering the concentration of compounds such as SDS (detergent) in the hybridisation buffer and/or exclusion of compounds such as dextran sulphate or polyethylene glycol (promoting molecular crowding) from the hybridisation buffer. Conventional hybridisation conditions are described in, for example, Sambrook (2001) Molecular Cloning: a laboratory manual, 3rd Edition Cold Spring Harbor Laboratory Press, CSH, New York, but the skilled craftsman will appreciate that numerous different hybridisation conditions may be designed in function of the known or the expected homology and/or length of the nucleic acid sequence. Sufficiently low stringency hybridisation conditions are particularly preferred (at least in the first instance) to isolate nucleic acids heterologous to the DNA sequences of the invention defined supra. An example of low stringency conditions is 4-6x SSC / 0.1-0.5% w/v SDS at 37-45°C for 2-3 hours. Depending on the source and concentration of the nucleic acid involved in the hybridisation, alternative conditions of stringency may be employed, such as medium stringency conditions. Examples of medium stringency conditions include 1-4x SSC / 0.25% w/v SDS at ≥ 45°C for 2-3 hours. An example of high stringency conditions includes 0.1 to 2x SSC / 0.1% w/v SDS at 60°C for 1-3 hours. The skilled man will be aware of various parameters which may be altered during hybridisation and washing and which will either maintain or change the stringency conditions. The stringency conditions may start low and be progressively increased until there is provided a hybridising nucleic acid, as defined hereinabove. Elements contributing to heterology include allelism, degeneration of the genetic code and differences in preferred codon usage.

Another variant 2xC2H2 zinc finger nucleic acid useful in practising the methods according to the present invention is an alternative splice variant of a nucleic acid sequence encoding a 2xC2H2 zinc finger protein. The term "alternative splice variant" as used herein encompasses variants of a nucleic acid sequence in which selected introns and/or exons have been excised, replaced or added. Such splice variants may be found in nature or may be manmade. Methods for making such splice variants are well known in the art. Splice variants suitable for use in the methods according to the invention may readily be determined for example by following the methods described in the Examples section by simply substituting the sequence used in the actual Example with the splice variant.

Another variant 2xC2H2 zinc finger nucleic acid useful in practising the methods according to the present invention is an allelic variant of a nucleic acid encoding a 2xC2H2 zinc finger protein. Allelic variants exist in nature and encompassed within the methods of the present invention is the use of these natural alleles. Allelic variants also encompass Single Nucleotide

Polymorphisms (SNPs), as well as Small Insertion/Deletion Polymorphisms (INDELS). The size of INDELS is usually less than 100 bp. SNPs and INDELS form the largest set of sequence variants in naturally occurring polymorphic strains of most organisms. Allelic variants suitable for use in the methods according to the invention may readily be determined for example by 5 following the methods described in the Examples section by simply substituting the sequence used in the actual Example with the allelic variant.

The present invention provides a method for modifying plant growth characteristics, comprising modifying expression in a plant of an alternative splice variant or expression in a plant of an 10 allelic variant of a nucleic acid encoding a 2xC2H2 zinc finger protein and/or by modifying level and/or activity in a plant of a 2xC2H2 zinc finger protein encoded by the alternative splice variant or allelic variant.

Examples of variant 2xC2H2 zinc finger proteins useful in practicing the methods of the 15 present invention are homologues, derivatives or functional fragments of a 2xC2H2 zinc finger protein.

“Homologues” of a 2xC2H2 zinc finger protein encompass peptides, oligopeptides, polypeptides, proteins and enzymes having amino acid substitutions, deletions and/or 20 insertions relative to the unmodified protein in question and having similar biological and functional activity as the unmodified protein from which they are derived. To produce such homologues, amino acids of the protein may be replaced by other amino acids having similar properties (such as similar hydrophobicity, hydrophilicity, antigenicity, propensity to form or break α -helical structures or β -sheet structures). Conservative substitution tables are well 25 known in the art (see for example Creighton (1984) Proteins. W.H. Freeman and Company). The homologues useful in the method according to the invention have at least in increasing order of preference 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 52%, 54%, 56%, 58%, 60%, 62%, 64%, 66%, 68%, 70%, 72%, 74%, 76%, 78%, 80%, 82%, 84%, 86%, 88%, 90%, 92%, 94%, 96%, 98% 30 sequence identity or similarity to an unmodified protein.

The percentage of identity may be calculated by using an alignment program well known in the art. For example, the percentage of identity may be calculated using the program GAP, or 35 needle (EMBOSS package) or stretcher (EMBOSS package) or the program align X, as a module of the vector NTI suite 5.5 software package, using the standard parameters (for example GAP penalty 5, GAP opening penalty 15, GAP extension penalty 6.6).

According to another embodiment of the present invention, the nucleic acid sequence useful in the methods of the present invention is a nucleic acid encoding a protein homologous to SEQ ID NO 2.

Methods for the search and identification of 2xC2H2 zinc finger protein homologues, for example STZ zinc finger homologues, would be well within the realm of a person skilled in the art. Such methods, involve screening sequence databases with the sequences provided by the present invention, for example SEQ ID NO 2 (or SEQ ID NO 1), preferably in a computer readable format. This sequence information may be available in public databases, that include but are not limited to Genbank (<http://www.ncbi.nlm.nih.gov/web/Genbank>), the European Molecular Biology Laboratory Nucleic acid Database (EMBL) (<http://www.ebi.ac.uk/ebi-docs/embl-db.html>) or versions thereof or the MIPS database (<http://mips.gsf.de/>). Different search algorithms and software for the alignment and comparison of sequences are well known in the art. Such methods include GAP, BESTFIT, BLAST, FASTA and TFASTA. GAP uses the algorithm of Needleman and Wunsch (J. Mol. Biol. 48: 443-453, 1970) to find the alignment of two complete sequences that maximises the number of matches and minimises the number of gaps. The BLAST algorithm calculates percent sequence identity and performs a statistical analysis of the similarity between the two sequences. The suite of programs referred to as BLAST programs has 5 different implementations: three designed for nucleotide sequence queries (BLASTN, BLASTX, and TBLASTX) and two designed for protein sequence queries (BLASTP and TBLASTN) (Coulson, Trends in Biotechnology: 76-80, 1994; Birren et al., GenomeAnalysis, 1: 543, 1997). The software for performing BLAST analysis is publicly available through the National Centre for Biotechnology Information.

Default blast parameters to find useful homologues of any of SEQ ID NO 1, SEQ ID NO 2 or any of SEQ ID NO 10 to SEQ ID NO 50, are, when comparing nucleotide sequence G (Cost to open a gap) 5, E (Cost to extend a gap default) 2, q (Penalty for a mismatch) -3, r (Reward for a match) 1, e (Expectation value (E)) 10.0, W (Word size) 11, V (Number of one-line descriptions) 100 and B (Number of alignments to show) 100. When comparing protein sequences, the default parameters are preferably G 11, E 1, e value 10.0, W 3, V 100 and B 100.

The above-mentioned analyses for comparing sequences, for the calculation of sequence identity and for the search for homologues, is preferentially done with full-length sequences or within a conserved region of the sequence. Therefore, these analyses may be based on a comparison of certain regions such as conserved domains, motifs or boxes.

The identification of such domains or motifs for examples the motif and boxes as represented by SEQ ID NO 5, 6, 7, 8 and 9, would also be well within the realm of a person skilled in the art and involves for example, a computer readable format of proteins of the present invention, the use of alignment software programs and the use of publicly available information on protein domains, conserved motifs and boxes. This protein domain information is available in the PRODOM (<http://www.biochem.ucl.ac.uk/bsm/dbbrowser/jj/prodomsrchjj.html>), PIR (<http://pir.georgetown.edu/>) or pFAM (<http://pfam.wustl.edu/>) database. For the identification of Zinc finger domains, such as the 2xC2H2 zinc finger domain, pFAM is preferred. Sequence analysis programs designed for motif searching may be used for identification of fragments, regions and conserved domains as mentioned above. Preferred computer programs would include but are not limited to MEME, SIGNALSCAN, and GENESCAN. A MEME algorithm (Version 3.0) may be found in the GCG package; or on the Internet site <http://www.sdsc.edu/MEME/meme>. SIGNALSCAN version 4.0 information is available on the Internet site <http://biosci.cbs.umn.edu/software/sigscan.html>. GENESCAN may be found on the Internet site <http://gnomic.stanford.edu/GENESCANW.html>.

At present, zinc finger motifs are subdivided in more than 40 different classes as can be found in the Pfam database of protein families present at the Sanger institute (<http://www.sanger.ac.uk/Software/Pfam/browse/Z.shtml>).

The C2H2 zinc finger (Zf-C2H2) motif is the classical zinc finger domain. It was first recognized in the transcription factor IIIA (TFIIIA) of Xenopus (Miller et al. 1985). The domain is typically 25 to 30 amino-acid residues in length. The following pattern describes the zinc finger *X-C-X(1-5)-C-X3-*X5-*X2-H-X(3-6)-[H/C] where X can be any amino acid, and numbers in brackets indicate the number of residues. The positions marked * are those that are important for the stable folding of the zinc finger. The final position can be either his or cys, while still being a C2H2 zinc finger domain. In view of recent publications on the design of zinc finger domains it becomes feasible also to replace one or more of the Cys or His amino acids, whilst still retaining the original functionality of the C2H2 domain. The residues separating the second Cys and the first His are mainly polar and basic. The canonical C2H2 zinc finger is composed of two short beta strands followed by an alpha helix. DNA binding of the zinc finger motif is mediated by amino terminal part of the alpha helix which binds the major groove in DNA binding zinc fingers. C2H2 domains have been shown to interact with RNA, DNA and proteins. The tetracoordination of a Zinc ion by the conserved cystein and histidine residues determines the conserved tertiary structure of the motif. Conserved hydrophobic residues are commonly found at positions -2 and also at 4 amino acids after the second cystein (that participates in zinc binding) and at position three before the first histidine (that participates in zinc binding). In

plant multi zinc finger proteins, spacing between the C2H2 domains is generally about 15 to about 65 amino acids.

Thus, plant zinc finger proteins are characterized by long spacers of diverse lengths between 5 adjacent fingers. Moreover, they are characterised by a highly conserved sequence of six amino acids, located within a putative DNA-contacting surface of each finger. Two forms of such conserved sequence are most commonly found in plant C2H2 zinc fingers, the QALGGH (SEQ ID NO 5) and the NNM/WQMH (SEQ ID NO 6). Despite the high sequence conservation of the QALGGH, some variants or the so-called 'modified type' occur in nature where one or 10 two amino acids can have a different form, most typically the +1 "Q" can be a "G", "K" or "R" (these amino acids share the same turn-like characteristic), the +2 "A" can be "S" (both of which share the characteristic of being small amino acids) or the +3 "L" can be "F" (these two amino acids are both hydrophobic). The QALGGH-motif as used herein comprises all these variants. In the NNM/WQMH motif at position 3 there is mostly an "M" or a "W".

15 Therefore, the present invention provides a method as described hereinabove, wherein said 2xC2H2 zinc finger protein comprises a QALGGH motif. Further, The present invention provides a s described hereinabove, wherein said 2xC2H2 zinc finger protein comprises a NNM/WQMH motif.

20 According to one embodiment of the invention, both C2H2 domains are of the same type. More preferably, both C2H2 zinc finger domains have the same conserved GALGGH or NNM/WQMH motif. According to another embodiment, each C2H2 zinc finger domain has a different conserved motif.

25 According to one embodiment, the 2xC2H2 protein useful in the methods of the present invention is characterized by an EAR motif, which is an ERF-Associated amphiphilic repression motif. This motif has been recognized in two unrelated types of transcription factors, namely the ERF transcription factors of the AP2 type and in the zinc finger transcription factors. In the latter class, the EAR motif is generally located at the C-terminus of the protein.

30 The pattern for the EAR motif has the conserved sequence hDLNh(X)P (SEQ ID NO 7), where "h" is a hydrophobic residue (any one of A,C,F,G,H,I,K,L,M,R,T,V,W,Y) most typically L/F/I and where "X" can be one (any amino acid) or no amino acid. A characteristic feature of the EAR motif is the alternation of hydrophilic and hydrophobic residues with the aspartic acid (D) residue being amphiphilic. Ohta et al. (The plant cell, 2001, 13, p1959-1968), which reference 35 is cited herein by reference, previously characterized EAR motifs present in 2xC2H2 zinc finger proteins.

Therefore, the present invention provides a method as described hereinabove, wherein the 2xC2H2 zinc finger protein comprises an EAR motif. According to one embodiment, the EAR motif is located in the C-terminal region of the protein, preferably between the second zinc finger domain and the C-terminus.

5

According to a further embodiment, the zinc finger proteins used in the methods of the present invention have two zinc finger domains and a nuclear localization signal (B-box). A cluster of basic amino acids that resembles the B-box (Basic box) were described by Chua et al. (EMBO 1992- 11, 241-9) and were hypothesized to be a nuclear localization signal for the protein.

10 These have been recognized in 2xC2H2 proteins (Sakamoto et al., Gene 248 (2000) 23-32). The cluster is rich in Lysine (K) and Arginine (R) residues. A consensus sequence defining the most frequent form of the B-box in 2xC2H2 genes is KR(S)KRXR (SEQ ID NO 8) where "S" at the 3rd position may be absent or present. However other variants may occur in nature that still retain the characteristic of being a charged region rich in basic amino acids. The location of 15 the basic box is most frequently at the N-terminus of the protein, but can also occur in other locations. It has been speculated that due to its basic nature the B-box could also participate in DNA binding.

Accordingly, the present invention provides a method as described hereinabove, wherein the 20 2xC2H2 zinc finger protein further comprises a B-box. According to one embodiment the B-box is located in the N-terminal region of the zinc finger protein. Preferably the proteins useful in the methods of the present invention have a B-box located between the N-terminus and the first zinc finger domain.

25 According to a further embodiment, the zinc finger proteins useful in the methods of the present invention have two C2H2 zinc finger domains and an L-box. A conserved motif, named L-box, of yet unknown function has been identified in 2xC2H2 proteins and has been described previously by Sakamoto et al. (Gene 248 (2000) 23-32). The L-box is typically located at the N-terminus, between the B-box and the first C2H2 zinc finger. The L-box is represented by the 30 sequence EXEXXAXCLXXL (SEQ ID NO 9). This region may be involved in protein-protein interactions. Zinc finger proteins lacking the L-box, may for example have serine rich regions at a similar position, which regions are putative sites for protein-protein interactions.

Therefore, the present invention provides a method as described hereinabove, wherein the 35 2xC2H2 protein comprises an L-box.

Particular zinc finger homologues useful in the methods of the present invention have one or more of the conserved motifs as depicted in SEQ ID NO 5, 6, 7, 8 and 9, or motifs that are 80% identical to these motifs or motifs that have conserved substitutions of amino acids. The 2xC2H2 protein as set forth in SEQ ID NO 2 comprises all the boxes as set forth in SEQ ID NO 5, 7, 8 and 9. All its paralogues and orthologues also comprise all of these boxes.

Homologues of a 2xC2H2 protein as presented in SEQ ID NO 2 and isolated from *Arabidopsis thaliana*, that are useful in the constructs and the methods of the present invention are also identified in other plant species.

Two special forms of homologue, orthologues and paralogues, are evolutionary concepts used to describe ancestral relationships of genes. The term "parologue" relates to a gene-duplication within the genome of a species leading to paralogous genes. The term "orthologue" relates to a homologous gene in different organisms due to ancestral relationship. The term "homologue" as used herein also encompasses paralogues and orthologues of the proteins useful in the methods according to the invention.

Othologues in other plant species may easily be found by performing a so-called reciprocal blast search. Orthologous genes can be identified by querying one or more gene databases with a query gene or protein of interest (SEQ ID NO 1 or 2), using for example BLAST program. The highest-ranking subject genes that result from the search are then again subjected to a BLAST analysis, and only those subject genes that match again with the query sequence (SEQ ID NO 1 or 2) are retained as true orthologous genes. For example, to find a rice orthologue of an *Arabidopsis thaliana* gene, one may perform a BLASTN or TBLASTX analysis on a rice database such as (but not limited to) the *Oryza sativa Nipponbare* database available at the NCBI website (<http://www.ncbi.nlm.nih.gov>) or the genomic sequences of rice (cultivars *indica* or *japonica*). In a next step, the obtained rice sequences are used in a reverse BLAST analysis using an *Arabidopsis* database. The results may be further refined when the resulting sequences are analysed with ClustalW and visualised in a neighbour joining tree. The method can be used to identify orthologues from many different species.

The closest homologues in other species (orthologues of the protein of SEQ ID NO 2), include those from a variety of dicot and monocot plants, for example from *Datisca glomerata* (AF119050_1, AAD26942, SEQ ID NO 10 and 11), from soybean (T09602, SCOF-1, SEQ ID NO 12 and 13), *Medicago sativa* (CAB77055.1, SEQ ID NO 14 and 15), from tobacco (T01985, SEQ ID NO 16 and 17) from rice, (AF332876_1, AAK01713.1, SEQ ID NO 18 and 19), from petunia (BAA05079.1, SEQ ID NO 20 and 21), from wheat (S39045 and BAA03901,

WZF1, SEQ ID NO 22 and 23), from *Capsicum annum* (SEQ ID NO 24 and 25), from turnip (T14408, T14409) and from sugarcane (CA279020).

Close homologues of the same species (paralogues of the protein of SEQ ID NO 2 from 5 *Arabidopsis thaliana*) are described below.

The MIPS database contains the sequence of the *Arabidopsis thaliana* genome with prediction and functional annotation of the proteins encoded. Searching this database with the STZ gene of SEQ ID NO 1 (MIPS accession number At1g27730), showed that in the *Arabidopsis* genome there are 2 genes encoding very close homologues of SEQ ID NO 2, At5g43170 10 (NM_123683, SEQ ID NO 32 and 33) and At5g04340 (NM_120516 SEQ ID NO 28 and 29), and 3 others with high similarity: At3g19580 (NM_112848, SEQ ID NO 26 and 27), At5g67450 (NM_126145, SEQ ID NO 34 and 35) and At3g49930 (NM_114853, SEQ ID NO 30-31). These genes are spread over 3 chromosomes, 1, 3 and 5. Similarly, a number of paralogues of the orthologue in Petunia have been isolated and sequenced. Advantageously, paralogues from 15 the same species may be used in the methods of the present invention.

Furthermore, a number of family members of the STZ protein of SEQ ID NO 2 have been found in *Arabisopsis*. The STZ gene and protein of SEQ ID NO 1 and 2 have been previously published in the database under the MIPS accession number At1g27730 or in Genbank under 20 the accession numbers NP_174094.1, X95573 or CAA64820. Additionally, several other cDNA's, isolated from other tissues or at different developmental stages of *Arabisopsis* have been reported and encode the same protein as that of SEQ ID NO 2. Such sequences 25 sequences deposited under the Genbank accession number AY034998, NM_102538, AC12375, X95573, AY063006, X98671, X98670, or AF250336. These isolates illustrate the differential expression of the STZ gene in different plant tissues at different developmental stages. The differential regulation of these different cDNA's is reflected by the differences at 30 their 5'UTR and the 3'UTR regions, while the encoded protein remains the same. Advantageously, the members of the same gene family as SEQ ID NO 1 or the members of the same family of any of the orthologues of SEQ ID NO 1, may be used in the methods of the present invention.

Other close homologues useful in the methods of the present invention are the sequences as deposited in the public database under the following accession numbers, which sequences are herein incorporated by reference: homologues isolated from *Petunia*: BAA21923.1, 35 BAA21922.1, BAA21926.1, BAA21925.1, BAA19110.1, BAA19926.1, BAA21924.1, BAA19111.1, BAA21921.1, BAA19114.1, BAA05076.1, BAA05079.1, CAA43111.1, BAA21920.1, BAA21919.1, BAA05077.1, BAA05078.1, BAA20137.1; homologues isolated

from *Arabidopsis*: CAA67229.1, BAC43454.1, NP_196054.1, AAM67193.1, NP_199131.1, NP_188592.1, NP_201546.1, NP_190562.1, NP_182037.1, BAC43008.1, Q8VWG3, CAC86393.1, CAC86168.1, CAC86167.1, CAC86166.1, CAB67667.1, CAC01747.1, CAB90936.1, CAB90935.1, CAB80245.1, CAB41188.1, CAA18741.1, CAA67234.1, 5 CAA67236.1, CAA67231.1, CAA67230.1, CAA67228.1, CAA67235.1, CAA67233.1, CAA67232.1, CAA67229.1, CAA64820.1 and homologues isolated from rice: BAB16855.1, AAO06972.1, CAC09475.1, BAB63718.1, P0683F02.21, BAB67885.1, P0031D11.19, BAB64114.1, AAK01713.1, AF332876_1, AAL76091.1, BAB67879.1, P0031D11.12 and BAC15513.1.

10

A phylogenetic tree may be constructed with all the homologues, paralogues and orthologues are defined herein above. Multiple alignment may be made using clustal W present in the VNTI (version 5.0) program with for example Gap opening penalty 10 and Gap extention 5. For making a phylogenetic tree the Phylic software package available at 15 <http://evolution.genetics.washington.edu/phylip.html> may be used. Sequences clustering around SEQ ID NO 1 or SEQ ID NO 2, Identify genes or proteins suitable for use in the methods of the present invention.

The sequence of SEQ ID NO 2 and its rice orthologue AF332876 (SEQ ID NO 19) have 36% 20 sequence identity when using the program Needle with the parameters Gap penalty 5 and Gap extension penalty 6. Therefore, homologues particularly useful in the methods of the present invention are homologues having 36% or more sequence identity with the 2xC2H2 zinc finger protein as presented in SEQ ID NO 2 or having 36% or more sequence identity to the closest orthologue of SEQ ID NO 2 from another species.

25

Preferred homologues useful in practicing the methods of the present invention are plant homologues, i.e. proteins obtained from a plant nucleic acid. More preferably, the nucleic acid sequence is from a dicot, more preferably from the family *Brassicaceae*, further preferably from *Arabidopsis thaliana*.

30

Preferably the 2xC2H2 zinc finger protein useful in the methods of the present invention belongs to the same gene family as the salt tolerant zinc finger protein (STZ) of *Arabidopsis thaliana*, or is a homologues thereof. The name ZAT10 can also be used to identify the STZ zinc finger protein of *Arabidopsis thaliana*.

35

Another variant of a zinc finger protein useful in the methods of the present invention is a substitutional variant. The term "Substitutional variants" of a protein refers to those variants in

which at least one residue in an amino acid sequence has been removed and a different residue inserted in its place. Amino acid substitutions are typically of single residues, but may be clustered depending upon functional constraints placed upon the polypeptide; insertions will usually be of the order of about 1-10 amino acid residues, and deletions will range from about

5 1-20 residues. Preferably, amino acid substitutions comprise conservative amino acid substitutions. Particular substitutional variants of the C2H2 zinc finger protein are substitutional variants in which one or more of the conserved Cys and/or His residues is replaced, whilst retaining the same zinc finger functionality. To retain the same functionality, the residues around these conserved Cys of His residues may also be substituted.

10

"Insertional variants" of a protein are those in which one or more amino acid residues are introduced into a predetermined site in said protein. Insertions can comprise amino-terminal and/or carboxy-terminal fusions as well as intra-sequence insertions of single or multiple amino acids. Generally, insertions within the amino acid sequence will be smaller than amino- or 15 carboxy-terminal fusions, of the order of about 1 to 10 residues. Examples of amino- or carboxy-terminal fusion proteins or peptides include the binding domain or activation domain of a transcriptional activator as used in the yeast two-hybrid system, phage coat proteins, (histidine)₆-tag, glutathione S-transferase-tag, protein A, maltose-binding protein, dihydrofolate reductase, Tag•100 epitope, c-myc epitope, FLAG[®]-epitope, lacZ, CMP (calmodulin-binding 20 peptide), HA epitope, protein C epitope and VSV epitope.

"Deletion variants" of a protein are characterised by the removal of one or more amino acids from the protein. Amino acid variants of a protein may readily be made using peptide synthetic techniques well known in the art, such as solid phase peptide synthesis and the like, or by 25 recombinant DNA manipulations. Methods for the manipulation of DNA sequences to produce substitution, insertion or deletion variants of a protein are well known in the art. For example, techniques for making substitution mutations at predetermined sites in DNA are well known to those skilled in the art and include M13 mutagenesis, T7-Gen *in vitro* mutagenesis (USB, Cleveland, OH), QuickChange Site Directed mutagenesis (Stratagene, San Diego, CA), PCR-mediated site-directed mutagenesis or other site-directed mutagenesis protocols.

The term "derivatives" refers to peptides, oligopeptides, polypeptides, proteins and enzymes which may comprise substitutions, deletions or additions of naturally and non-naturally occurring amino acid residues compared to the amino acid sequence of a naturally-occurring 35 form of the 2xC2H2 protein such as for example the 2xC2H2 zinc finger protein as presented in SEQ ID NO 2. "Derivatives" of a 2xC2H2 zinc finger protein encompass peptides, oligopeptides, polypeptides, proteins and enzymes which may comprise naturally occurring

altered, glycosylated, acylated or non-naturally occurring amino acid residues compared to the amino acid sequence of a naturally-occurring form of the polypeptide. A derivative may also comprise one or more non-amino acid substituents compared to the amino acid sequence from which it is derived, for example a reporter molecule or other ligand, covalently or non-
5 covalently bound to the amino acid sequence such as, for example, a reporter molecule which is bound to facilitate its detection, and non-naturally occurring amino acid residues relative to the amino acid sequence of a naturally-occurring protein.

Another variant of a 2xC2H2 zinc finger protein useful in the methods of the present invention
10 is an active fragment of a zinc finger protein. "Active fragments" of a 2xC2H2 zinc finger protein encompasses at least five contiguous amino acid residues of a protein, which residues retain similar biological and/or functional activity to the naturally occurring protein. For example, useful fragments comprise at least 10 contiguous amino acid residues of a 2xC2H2 zinc finger protein. Other preferred fragments are fragments of a 2xC2H2 zinc finger protein
15 starting at the second or third or further internal methionine residues. These fragments originate from protein translation, starting at internal ATG codons. Functional fragments of a 2xC2H2 zinc finger protein useful in practising the methods of the present invention may have one, two or no C2H2 domains, without affecting its functionality in the methods of the present invention.

20 According to a preferred feature of the present invention, enhanced or increased expression of a nucleic acid encoding a 2xC2H2 zinc finger protein is envisaged. Methods for obtaining enhanced or increased expression of genes or gene products are well documented in the art and include, for example, over-expression driven by a strong promoter, the use of transcription enhancers or translation enhancers. The term over-expression as used herein means any form
25 of expression that is additional to the original wild-type expression level. Preferably the nucleic acid to be introduced into the plant and/or the nucleic acid that is to be overexpressed in the plant is in the sense direction with respect to the promoter to which it is operably linked. Preferably, the nucleic acid sequence represented by SED ID NO 1 is over-expressed in a plant. However, it should be clear that the applicability of the invention is not limited to use of
30 the nucleic acid represented by SEQ ID NO 1 nor to the nucleic acid sequence encoding the amino acid sequence of SEQ ID NO 2, but that other nucleic acid sequences encoding homologues, derivatives or active fragments of SED ID NO 1 or SED ID NO 2 may be useful in the methods of the present invention. Examples of nucleic acids or proteins are provided in SEQ ID NO 10 to SEQ ID NO 50.

35 Alternatively and/or additionally, increased expression of a 2xC2H2 encoding gene or increased level and/or activity of a 2xC2H2 protein in a plant cell, is achieved by mutagenesis.

For example these mutations may be responsible for altered control of the 2xC2H2 gene, resulting in more expression of the gene, relative to the wild-type gene. Mutations can also cause conformational changes in a protein, resulting in more activity and/or higher levels of the 2xC2H2 protein.

5

Modifying gene expression (whether by a direct or indirect approach) encompasses altered transcript levels of a gene. Altered transcript levels may be sufficient to induce certain phenotypic effects, for example via the mechanism of cosuppression. Here the overall effect of introduction of a transgene is that there is less activity in the cell of the protein encoded by a native gene having homology to the introduced transgene. Therefore, according to another embodiment of the present invention, there is provided a method for modifying growth characteristics in a plant, comprising decreasing expression of a gene encoding a 2xC2H2 zinc finger protein or decreasing level and/or activity of a 2xC2H2 zinc finger protein. Examples of decreasing expression, level and/or activity of a protein in a cell are well documented in the art and include, for example, downregulation of expression by anti-sense techniques, RNAi techniques, small interference RNAs (siRNAs) and microRNA (miRNA).

Another method for downregulation of gene expression or gene silencing comprises use of ribozymes, for example as described in Atkins et al. 1994 (WO 94/00012), Lenee et al. 1995 (WO 95/03404), Lutziger et al. 2000 (WO 00/00619), Prinsen et al. 1997 (WO 97/3865) and Scott et al. 1997 (WO 97/38116).

Gene silencing may also be achieved by insertion mutagenesis (for example, T-DNA insertion or transposon insertion) or by gene silencing strategies as described by, among others, Angell and Baulcombe 1998 (WO 98/36083), Lowe et al. 1989 (WO 98/53083), Lederer et al. 1999 (WO 99/15682) or Wang et al. 1999 (WO 99/53050).

Expression of an endogenous gene may also be reduced if it contains a mutation. Such a mutation or such a mutant gene may be isolated and introduced into the same or different plant species in order to obtain plants having modified growth characteristics. Examples of such mutants are dominant negative mutants of a 2xC2H2 zinc finger gene.

Genetic constructs aimed at silencing gene expression may comprise the 2xC2H2 zinc finger nucleic acid, for example as represented by SEQ ID NO 1 (or one or more portions thereof or a sequence capable of hybridising therewith), in a sense and/or antisense orientation relative to the promoter sequence. The sense or antisense copies of at least part of the endogenous gene in the form of direct or inverted repeats may also be utilised in the methods according to

the invention. The growth characteristics of plants may also be modified by introducing into a plant at least part of an antisense version of the nucleotide sequence represented by SEQ ID NO 1.

5 According to a further embodiment of the present invention, genetic constructs and vectors to facilitate introduction and/or to facilitate expression of the 2xC2H2 zinc finger nucleotide sequences useful in the methods according to the invention are provided. Therefore, according to the present invention, there is provided a construct comprising:

- (i) a nucleic acid capable of modifying expression of a nucleic acid encoding a 10 2xC2H2 zinc finger protein and/or modifying level and/or activity of a 2xC2H2 zinc finger protein;
- (ii) one or more control sequence capable of driving expression of the nucleic acid sequence of (i); and optionally
- (iii) a transcription termination sequence.

15 Constructs useful in the methods according to the present invention may be constructed using recombinant DNA technology well known to persons skilled in the art. The gene constructs may be inserted into vectors, which may be commercially available, suitable for transforming into plants and suitable for expression of the gene of interest in the transformed cells.

20 Preferably the genetic construct is a plant expression vector.

The nucleic acid according to (i) is advantageously any of the nucleic acids described hereinbefore. A preferred nucleic acid is the nucleic acid represented by SEQ ID NO 1 or a variant thereof as hereinbefore defined, or is a nucleic acid sequence encoding a sequence 25 represented by SEQ ID NO 2 or a variant as hereinbefore defined. For example such variants encode a protein as presented in any of SEQ ID NO 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 39, 42, 44, 46, 48 and 50.

30 The terms "regulatory element" and "control sequence" are used herein interchangeably and are to be taken in a broad context to refer to regulatory nucleic acids capable of effecting expression of the sequences to which they are operably linked. Encompassed by the aforementioned terms are promoters. A "promoter" encompasses transcriptional regulatory sequences derived from a classical eukaryotic genomic gene (including the TATA box which is required for accurate transcription initiation, with or without a CCAAT box sequence) and 35 additional regulatory elements (i.e. upstream activating sequences, enhancers and silencers) which alter gene expression in response to developmental and/or external stimuli, or in a tissue-specific manner. Also included within the term is a transcriptional regulatory sequence

of a classical prokaryotic gene, in which case it may include a -35 box sequence and/or -10 box transcriptional regulatory sequences. The term "regulatory element" also encompasses a synthetic fusion molecule or derivative which confers, activates or enhances expression of a nucleic acid molecule in a cell, tissue or organ. The term "operably linked" as used herein 5 refers to a functional linkage between the promoter sequence and the gene of interest, such that the promoter sequence is able to initiate transcription of the gene of interest. Preferably, the gene of interest is operably linked to a promoter in a sense direction.

Advantageously, any type of promoter may be used to drive expression of the nucleic acid 10 sequence depending on the desired outcome.

Promoters useful for the present invention are described in EP 03075331.3, which promoters and sequences are incorporated herein by reference.

Other examples of preferred promoters are presented in Table I (a) to (c), which promoters or 15 derivatives thereof are useful in the methods and/or in making the constructs of the present invention. Accordingly, genetic constructs comprising of the nucleic acids of (i), for example a 2xC2H2 nucleic acid, and at least part of a promoter from Table I (a) to (c) or from EP 03075331.3, preferably, wherein said parts are operably linked, are also provided by the present invention.

20 According to another embodiment, the nucleic acid of (i) is operably linked to a constitutive promoter. The term "constitutive" as defined herein refers to a promoter that is expressed substantially continuously. Furthermore, preferably the constitutive promoter is a ubiquitous promoter, which is expressed in more than one, preferably in most or substantially all tissues 25 of the plant. Preferably, the constitutive promoter to be used in the methods of the present invention, or cloned in the genetic constructs of the present invention, is a plant promoter, preferably a constitutive promoter, such as a GOS2 promoter or a promoter with similar strength and/or similar expression pattern. Preferably plant promoters derived from a plant nucleic acid are used. Alternatively, promoters operable in plant, such as promoters derived 30 from plant pathogens are used.

According to another embodiment of the invention, the nucleic acid of (i) is operably linked to a plant promoter, preferably a tissue-preferred promoter. The term "tissue-preferred" as used herein refers to a promoter that is expressed predominantly in at least one tissue or organ. For 35 example, the tissue-preferred promoter is a seed-preferred promoter, such as a pWS18 (Joshee et al. Plant Cell Physiol. 1998 Jan;39(1):64-72.) or a promoter of similar strength and/or similar expression pattern.

Promoters with similar strength and/or similar expression pattern may be found by coupling the promoter to a reporter gene and checking the function of the reporter gene in different tissues of a plant. One suitable reporter gene is beta-glucuronidase and the colorimetric GUS staining 5 to visualize the beta-glucuronidase activity in a plant tissue is well known to a person skilled in the art.

Table I (a): flower preferred promoters useful in the present invention. Sequences of these 10 promoters are described in the cited reference, which sequences are herein incorporated by reference.

Gene	Expression	Reference
AtPRP4	flowers	http://salus.medium.edu/mmg/tierney/html
chalone synthase (chsA)	flowers	Van der Meer, et al., <i>Plant Mol. Biol.</i> 15, 95-109, 1990.
LAT52	anther	Twell et al <i>Mol. Gen Genet.</i> 217:240-245 (1989)
apetala-3	flowers	

Table I (b): seed-preferred promoters useful in the present invention. Sequences of these 15 promoters are described in the cited reference, which sequences are herein incorporated by reference.

Gene	Expression	Reference
seed-specific genes	seed	Simon, et al., <i>Plant Mol. Biol.</i> 5: 191, 1985; Scofield, et al., <i>J. Biol. Chem.</i> 262: 12202, 1987.; Baszczynski, et al., <i>Plant Mol. Biol.</i> 14: 633, 1990.
Brazil Nut albumin	seed	Pearson, et al., <i>Plant Mol. Biol.</i> 18: 235-245, 1992.
legumin	seed	Ellis, et al., <i>Plant Mol. Biol.</i> 10: 203-214, 1988.
glutelin (rice)	seed	Takaiwa, et al., <i>Mol. Gen. Genet.</i> 208: 15-22, 1986; Takaiwa, et al., <i>FEBS Letts.</i> 221: 43-47, 1987.
zein	seed	Matzke et al <i>Plant Mol Biol.</i> 14(3):323-32 1990
napA	seed	Stalberg, et al, <i>Planta</i> 199: 515-519, 1996.

wheat LMW and HMW glutenin-1	endosperm	Mol Gen Genet 216:81-90, 1989; NAR 17:461-2, 1989
wheat SPA	seed	Albani <i>et al</i> , Plant Cell, 9: 171-184, 1997
wheat α , β , γ -gliadins	endosperm	EMBO 3:1409-15, 1984
barley <i>ltr1</i> promoter	endosperm	
barley B1, C, D, hordelin	endosperm	Theor Appl Gen 98:1253-62, 1999; Plant J 4:343-55, 1993; Mol Gen Genet 250:750-60, 1996
barley DOF	endosperm	Mena <i>et al</i> , The Plant Journal, 116(1): 53-62, 1998
<i>blz2</i>	endosperm	EP99106056.7
synthetic promoter	endosperm	Vicente-Carbachosa <i>et al</i> , Plant J. 13: 629 - 640, 1998.
rice prolamin NRP33	endosperm	Wu <i>et al</i> , Plant Cell Physiology 39(8) 885-889, 1998
rice α -globulin Glb-1	endosperm	Wu <i>et al</i> , Plant Cell Physiology 39(8) 885-889, 1998
rice OSH1	embryo	Sato <i>et al</i> , Proc. Natl. Acad. Sci. USA, 93: 8117-8122, 1996
rice α -globulin REB/OHP-1	endosperm	Nakase <i>et al</i> . Plant Mol. Biol. 33: 513-522, 1997
rice ADP-glucose PP	endosperm	Trans Res 6:157-68, 1997
maize ESR gene family	endosperm	Plant J 12:235-46, 1997
sorghum γ -kafirin	endosperm	PMB 32:1029-35, 1996
KNOX	embryo	Postma-Haarsma <i>et al</i> , Plant Mol. Biol. 39:257-71, 1999
rice oleosin	embryo and aleuron	Wu <i>et al</i> , J. Biochem., 123:386, 1998
sunflower oleosin	seed (embryo and dry seed)	Cummins, <i>et al.</i> , Plant Mol. Biol. 19: 873-876, 1992

Table I (c): constitutive promoters useful in the present invention. Sequences of these promoters are described in the cited reference, which sequences are herein incorporated by reference.

Gene	Expression	Reference
Actin	constitutive	McElroy <i>et al</i> , Plant Cell, 2: 163-171, 1990
CAMV 35S	constitutive	Odell <i>et al</i> , Nature, 313: 810-812,

		1985
CaMV 19S	constitutive	Nilsson <i>et al.</i> , <i>Physiol. Plant.</i> 100:456-462, 1997
GOS2	constitutive	de Pater <i>et al.</i> , <i>Plant J</i> Nov;2(6):837-44, 1992
ubiquitin	constitutive	Christensen <i>et al.</i> , <i>Plant Mol. Biol.</i> 18: 675-689, 1992
rice cyclophilin	constitutive	Buchholz <i>et al.</i> , <i>Plant Mol Biol.</i> 25(5): 837-43, 1994
maize H3 histone	constitutive	Lepetit <i>et al.</i> , <i>Mol. Gen. Genet.</i> 231:276-285, 1992
actin 2	constitutive	An <i>et al.</i> , <i>Plant J.</i> 10(1); 107-121, 1996

Optionally, one or more terminator sequences may also be used in the construct introduced into a plant. The term "terminator" encompasses a control sequence which is a DNA sequence at the end of a transcriptional unit which signals 3' processing and polyadenylation of a primary

5 transcript and termination of transcription. Additional regulatory elements may include transcriptional as well as translational enhancers. Those skilled in the art will be aware of terminator and enhancer sequences which may be suitable for use in the invention. Such sequences would be known or may readily be obtained by a person skilled in the art.

10 The genetic constructs of the invention may further include an origin of replication sequence which is required for maintenance and/or replication in a specific cell type. One example is when a genetic construct is required to be maintained in a bacterial cell as an episomal genetic element (e.g. plasmid or cosmid molecule). Preferred origins of replication include, but are not limited to, the f1-ori and colE1.

15 The genetic construct may optionally comprise a selectable marker gene. As used herein, the term "selectable marker gene" includes any gene which confers a phenotype on a cell in which it is expressed to facilitate the identification and/or selection of cells which are transfected or transformed with a genetic construct of the invention. Suitable markers may be selected from 20 markers that confer antibiotic or herbicide resistance. Cells containing the recombinant DNA will thus be able to survive in the presence of antibiotic or herbicide concentrations that kill untransformed cells. Examples of selectable marker genes include genes conferring resistance to antibiotics (such as nptII encoding neomycin phosphotransferase capable of phosphorylating neomycin and kanamycin, or hpt encoding hygromycin phosphotransferase

capable of phosphorylating hygromycin), to herbicides (for example bar which provides resistance to Basta; aroA or gox providing resistance against glyphosate), or genes that provide a metabolic trait (such as manA that allows plants to use mannose as sole carbon source). Visual marker genes result in the formation of colour (for example beta-glucuronidase,

5 GUS), luminescence (such as luciferase) or fluorescence (Green Fluorescent Protein, GFP, and derivatives thereof). Further examples of suitable selectable marker genes include the ampicillin resistance (Ampr), tetracycline resistance gene (Tcr), bacterial kanamycin resistance gene (Kanr), phosphinothricin resistance gene, and the chloramphenicol acetyltransferase (CAT) gene, amongst others

10

The present invention also encompasses plants obtainable by the methods according to the present invention. The present invention therefore provides plants obtainable by the method according to the present invention, which plants have modified growth characteristics, which plants have altered 2xC2H2 zinc finger protein level and/or activity and/or altered expression of 15 a nucleic acid sequence encoding a 2xC2H2 zinc finger protein.

Therefore, according to one aspect of the present invention, there is provided a method for the production of plants, having modified growth characteristics, comprising introducing, into a plant, a nucleic acid capable of modifying activity of a 2xC2H2 zinc finger protein and/or 20 capable of modifying expression of a 2xC2H2 zinc-finger gene. According to a further embodiment of the present invention, there is provided a method for the production of transgenic plants having modified growth characteristics, comprising introduction and expression in a plant of a 2xC2H2 nucleic acid.

25 More specifically, the present invention provides a method for the production of transgenic plants having modified growth characteristics, which method comprises:

- (i) introducing into a plant or plant cell a 2xC2H2 zinc finger nucleic acid;
- (iii) cultivating the plant cell under conditions promoting plant growth.

30 The growth characteristic may be any of the characteristics defined hereinunder.

The 2xC2H2 zinc finger nucleic acid includes all variant nucleic acids as described herein before and includes all nucleic acids encoding all variant proteins as described herein before.

35 Cultivating the plant cell under conditions promoting plant growth, may or may not include regeneration and or growth to maturity.

The protein itself and/or the nucleic acid itself may be introduced directly into a plant cell or into the plant itself (including introduction into a tissue, organ or any other part of the plant). According to a preferred feature of the present invention, the nucleic acid is preferably introduced into a plant by transformation.

5

The term "transformation" as referred to herein encompasses the transfer of an exogenous polynucleotide into a host cell, irrespective of the method used for transfer. Plant tissue capable of subsequent clonal propagation, whether by organogenesis or embryogenesis, may be transformed with a genetic construct of the present invention and a whole plant regenerated therefrom. The particular tissue chosen will vary depending on the clonal propagation systems available for, and best suited to, the particular species being transformed. Exemplary tissue targets include leaf disks, pollen, embryos, cotyledons, hypocotyls, megagametophytes, callus tissue, existing meristematic tissue (e.g., apical meristem, axillary buds, and root meristems), and induced meristem tissue (e.g., cotyledon meristem and hypocotyl meristem). The polynucleotide may be transiently or stably introduced into a host cell and may be maintained non-integrated, for example, as a plasmid. Alternatively, it may be integrated into the host genome. The resulting transformed plant cell can then be used to regenerate a transformed plant in a manner known to persons skilled in the art.

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Transformation of a plant species is now a fairly routine technique. Advantageously, any of several transformation methods may be used to introduce the nucleic acid of interest (e.g. the 2xC2H2 nucleic acid) into a suitable ancestor cell. Transformation methods include the use of liposomes, electroporation, chemicals that increase free DNA uptake, injection of the DNA directly into the plant, particle gun bombardment, transformation using viruses or pollen and microprojection. Methods may be selected from the calcium/polyethylene glycol method for protoplasts (Krens, F.A. et al., 1982, *Nature* 296, 72-74; Negruțiu I. et al., June 1987, *Plant Mol. Biol.* 8, 363-373); electroporation of protoplasts (Shillito R.D. et al., 1985 *Bio/Technol* 3, 1099-1102); microinjection into plant material (Crossway A. et al., 1986, *Mol. Gen Genet* 202, 179-185); DNA or RNA-coated particle bombardment (Klein T.M. et al., 1987, *Nature* 327, 70) infection with (non-integrative) viruses and the like. A preferred transformation method is an *Agrobacterium* mediated transformation method.

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Transgenic rice plants expressing a 2xC2H2 gene are preferably produced via *Agrobacterium*-mediated transformation using any of the well-known methods for rice transformation, such as the ones described in any of the following: published European patent application EP 1198985 A1, Aldemita and Hodges (*Planta*, 199, 612-617, 1996); Chan et al. (*Plant Mol. Biol.* 22 (3) 491-506, 1993); Hiei et al. (*Plant J.* 6 (2) 271-282, 1994); which disclosures are incorporated

by reference herein as if fully set forth. In the case of corn transformation, the preferred method is as described in either Ishida et al. (Nat. Biotechnol. 1996 Jun; 14(6): 745-50) or Frame et al. (Plant Physiol. 2002 May; 129(1): 13-22), which disclosures are incorporated by reference herein as if fully set forth.

5

Generally after transformation, plant cells or cell groupings are selected for the presence of one or more markers which are encoded by plant-expressible genes co-transferred with the gene of interest, following which the transformed material is regenerated into a whole plant.

10 Following DNA transfer and regeneration, putatively transformed plants may be evaluated, for instance using Southern analysis, for the presence of the gene of interest, copy number and/or genomic organisation. Alternatively or additionally, expression levels of the newly introduced DNA may be undertaken using Northern and/or Western analysis, both techniques being well known to persons having ordinary skill in the art.

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The generated transformed plants may be propagated by a variety of means, such as by clonal propagation or classical breeding techniques. For example, a first generation (or T1) transformed plant may be selfed to give homozygous second generation (or T2) transformants, and the T2 plants further propagated through classical breeding techniques.

20

The generated transformed organisms may take a variety of forms. For example, they may be chimeras of transformed cells and non-transformed cells; clonal transformants (e.g., all cells transformed to contain the expression cassette); grafts of transformed and untransformed tissues (e.g., in plants, a transformed rootstock grafted to an untransformed scion).

25

The present invention clearly extends to any plant cell or plant produced by any of the methods described herein, and to all plant parts and propagules thereof. The present invention extends further to encompass the progeny of a primary transformed or transfected cell, tissue, organ or whole plant that has been produced by any of the aforementioned methods, the only

30 requirement being that progeny exhibit the same genotypic and/or phenotypic characteristic(s) as those produced in the parent by the methods according to the invention. The invention also includes host cells having modified expression and/or level and/or activity of a 2xC2H2 zinc finger protein. Such host cells for example comprise genetic constructs as mentioned above. Preferred host cells according to the invention are derived from a plant, algae, bacterium, 35 fungus, yeast, insect or animal. The invention also extends to harvestable parts of a plant such as but not limited to seeds, leaves, fruits, flowers, petals, stamen, stem cultures, stem, rhizomes, roots, tubers, bulbs or cotton fibers.

The term "plant" as used herein encompasses whole plants, ancestors and progeny of the plants and plant parts, including seeds, shoots, stems, roots (including tubers), and plant cells, tissues and organs. The term "plant" also encompasses suspension cultures, embryos, 5 meristematic regions, callus tissue, leaves, gametophytes, sporophytes, pollen, and microspores. Plants that are particularly useful in the methods of the invention include all plants which belong to the superfamily *Viridiplanteae*, in particular monocotyledonous and dicotyledonous plants including, fodder or forage legumes, ornamental plants, food crop, tree, or shrub selected from the list comprising *Acacia spp.*, *Acer spp.*, *Actinidia spp.*, *Aesculus spp.*, 10 *Agathis australis*, *Albizia amara*, *Alsophila tricolor*, *Andropogon spp.*, *Arachis spp.*, *Areca catechu*, *Astelia fragrans*, *Astragalus cicer*, *Balkiaea plurijuga*, *Betula spp.*, *Brassica spp.*, *Bruguiera gymnorhiza*, *Burkea africana*, *Butea frondosa*, *Cadaba farinosa*, *Calliandra spp.*, *Camellia sinensis*, *Canna indica*, *Capsicum spp.*, *Cassia spp.*, *Centroema pubescens*, *Chaenomeles spp.*, *Cinnamomum cassia*, *Coffea arabica*, *Colophospermum mopane*, 15 *Coronilla varia*, *Cotoneaster serotina*, *Crataegus spp.*, *Cucumis spp.*, *Cupressus spp.*, *Cyathea dealbata*, *Cydonia oblonga*, *Cryptomeria japonica*, *Cymbopogon spp.*, *Cynthea dealbata*, *Cydonia oblonga*, *Dalbergia monetaria*, *Davallia divaricata*, *Desmodium spp.*, *Dicksonia squarosa*, *Diheteropogon amplectens*, *Dioclea spp.*, *Dolichos spp.*, *Dorycnium rectum*, *Echinochloa pyramidalis*, *Ehrartia spp.*, *Eleusine coracana*, *Eragrostis spp.*, *Erythrina spp.*, *Eucalyptus spp.*, *Euclea schimperi*, *Eulalia villosa*, *Fagopyrum spp.*, *Feijoa sellowiana*, 20 *Fragaria spp.*, *Flemingia spp.*, *Freycinetia banksii*, *Geranium thunbergii*, *Ginkgo biloba*, *Glycine javanica*, *Gliricidia spp.*, *Gossypium hirsutum*, *Grevillea spp.*, *Guibourtia coleosperma*, *Hedysarum spp.*, *Hemarthria altissima*, *Heteropogon contortus*, *Hordeum vulgare*, *Hyparrhenia rufa*, *Hypericum erectum*, *Hyperthelia dissoluta*, *Indigo incarnata*, *Iris spp.*, *Leptarrhena pyrolifolia*, *Lespediza spp.*, *Lettuca spp.*, *Leucaena leucocephala*, *Loudetia simplex*, *Lotus bainesii*, *Lotus spp.*, *Macrotyloma axillare*, *Malus spp.*, *Manihot esculenta*, *Medicago sativa*, *Metasequoia glyptostroboides*, *Musa sapientum*, *Nicotianum spp.*, *Onobrychis spp.*, *Ornithopus spp.*, *Oryza spp.*, *Peltophorum africanum*, *Pennisetum spp.*, *Persea gratissima*, *Petunia spp.*, *Phaseolus spp.*, *Phoenix canariensis*, *Phormium cookianum*, *Photinia spp.*, 25 *Picea glauca*, *Pinus spp.*, *Pisum sativum*, *Podocarpus totara*, *Polygonarthria fleckii*, *Polygonarthria squarrosa*, *Populus spp.*, *Prosopis cineraria*, *Pseudotsuga menziesii*, *Pterolobium stellatum*, *Pyrus communis*, *Quercus spp.*, *Rhaphiolepis umbellata*, *Rhopalostylis sapida*, *Rhus natalensis*, *Ribes grossularia*, *Ribes spp.*, *Robinia pseudoacacia*, *Rosa spp.*, *Rubus spp.*, *Salix spp.*, *Schyzachyrium sanguineum*, *Sciadopitys verticillata*, *Sequoia sempervirens*, *Sequoiadendron giganteum*, *Sorghum bicolor*, *Spinacia spp.*, *Sporobolus fimbriatus*, *Stiburus alopecuroides*, *Stylosanthos humilis*, *Tadehagi spp.*, *Taxodium distichum*, *Themeda triandra*, *Trifolium spp.*, *Triticum spp.*, *Tsuga heterophylla*, *Vaccinium*

spp., *Vicia spp.* *Vitis vinifera*, *Watsonia pyramidata*, *Zantedeschia aethiopica*, *Zea mays*, amaranth, artichoke, asparagus, broccoli, brussel sprout, cabbage, canola, carrot, cauliflower, celery, collard greens, flax, kale, lentil, oilseed rape, okra, onion, potato, rice, soybean, straw, sugarbeet, sugar cane, sunflower, tomato, squash, and tea, trees and algae amongst others.

5 According to a preferred embodiment of the present invention, the plant is a crop plant such soybean, sunflower, canola, alfalfa, rapeseed, cotton, tomato, potato or tobacco. According to another preferred embodiment of the present invention, the plant is a monocotyledonous plant, such as sugar cane, further preferably a cereal, most preferably the plant is selected from the group consisting of rice, maize, wheat, barley, millet, rye or oats.

10 In a particular embodiment of the present invention, proteins of one plant species (for example *Arabidopsis*) are introduced in another plant species (for example rice). It has been shown in the present invention that plant growth characteristics are improved by introduction of a 2xC2H2 zinc finger gene or protein from a dicot into a monocot.

15 According to a particular embodiment of the invention, there are provided methods as described above, wherein the plant is a monocot. More preferably the plant is rice or corn.

20 Advantageously, performance of the methods according to the present invention leads to plants having modified growth characteristics.

The term "growth characteristic" as used herein, preferably refers to anyone or more of, but is not limited to, yield, architecture and cycle time.

25 The term "yield" means the amount of harvested material. For crop plants yield also means the amount of harvested material per acre of production. Depending on the crop the harvested part of the plant may be a different part or tissue of the plant, such as seed (e.g. rice, sorghum or corn when grown for seed); total above-ground biomass (e.g. for corn, when used as silage), root (e.g. sugarbeet), fruit (e.g. tomato), cotton fibers, or any other part of the plant which is of economic value. "Yield" also encompasses yield stability of the plants, meaning that year after year, one can obtain the same yield from the progeny of the plants, without too much interference of external factors, such as weather conditions. "Yield" also encompasses yield potential, which as the maximum obtainable yield.

30 Yield maybe dependent on a number of yield components. The parameters for these components are well known by a person skilled in the art. For example breeders are well aware of the specific yield components and the corresponding parameters for the crop they are aiming to improve.

For example key yield components for corn include number of plants per hectare or acre, number of ears per plant, number of rows (of seeds) per ear, number of kernels per row, and thousand kernel weight. For silage corn typical parameters are the above ground biomass and energy content.

5 Key yield components for rice include number of plants per hectare or acre, number of panicles per plant, number of spikelets per panicle, seed filling rate (number of filled seeds) and thousand kernel weight. Preferentially methods for increasing yield of rice encompass increased number of flowers per panicle and an increased number of filled seeds. The parameter of increased total number of seeds may be linked to increased number of flowers.

10 "Yield" further encompasses typical biomass components, such as above ground parts of a plant and the root system. General biomass parameters are area and dry weight. Specific parameters for above ground biomass further encompass above ground area and plant height. Specific parameters for the root system encompass root ratio, root length and penetration depth, root branching, root hair density, root pulling resistance and aerenchyma formation.

15 The plants of the present invention are characterized by increased number of filled seeds, increased total seed weight, increased total number of seeds and increased harvest index. Therefore the methods of the present invention are particularly favorable to be applied in cereals such as rice and corn (maize). Accordingly, a particular embodiment of the present

20 invention relates to a method to increase yield of corn, comprising modifying expression of a nucleic acid encoding a 2xC2H2 zinc finger protein.

25 The plants of the present invention are characterized by an increase in thousand kernel weight and therefore the seed size or seed volume and/or the seed content and/or seed composition are altered by the methods of the present invention. The seeds provided by the methods of the present invention may have more nutritional value, more starch and/or more oil, possibly due to their bigger size.

30 The plants of the present invention are characterized by more above ground area. Therefore, the methods of the present invention are particularly favorable for crops grown for their green tissue and/or grown for their above ground biomass. The methods of the present invention are particularly useful for grasses, forage crops (such as forage corn (maize), clover, medicago etc.), trees, sugar cane etc.

35 The improvement in yield as obtained by the methods of the invention, may be obtained as a result of improvement of one or more of the above mentioned yield components and/or parameters.

The term "architecture" as used herein encompasses the appearance or morphology of a plant, including any one or more structural features or combination of structural features thereof. Such structural features include the shape, size, number, position, texture, 5 arrangement, and pattern of any cell, tissue or organ or groups of cells, tissues or organs of a plant, including the root, leaf, shoot, stem, petiole, trichome, flower, petal, stigma, style, stamen, pollen, ovule, seed, embryo, endosperm, seed coat, aleurone, fibre, cambium, wood, heartwood, parenchyma, aerenchyma, sieve element, phloem or vascular tissue, amongst others. Particular architectural characteristics that may be modified by the methods of the 10 present invention are increased plant height, increased number or size of stems or stalks or tillers or panicles or pedicles, increased number or size of inflorescences, increased branching of for example of tassels and ears or altered flowering characteristics. A preferred architectural characteristic that may be modified by the methods of the present invention is leaf architecture. The term "leaf architecture" as used herein comprises typical leaf characteristics such as 15 length, width, thickness, cell number, cell size and greenness.

Typically, the plants of the present invention display increased leaf surface area and /or increased leaf blade width. This trait is particularly important as it allows the plant to optimize the shape of its leaf to maximize the area used for photosynthesis. For that purpose, preferably 20 the leaf blade is widened, but alternatively, the leaves are longer or smaller or rounder. These effects may lead to more healthy plants. Alternatively, this trait attributes aesthetic properties to the plant such as greenness and stronger leafs.

"cycle time" of the plant as used herein means the time wherein a plant reaches 90% of its 25 maximum total area. This parameter is an indication of the duration of the vegetative growth. Prolonged vegetative growth was only displayed in some of the plants according to the present invention and may be controlled by choice of the transformation event and/or by choice of the promoter driving the 2xC2H2 nucleic acid. For example this characteristic was not displayed when a seed-preferred promoter was used.

30 Other "growth characteristics" that may be improved by the methods of the present invention are growth rate, early vigour, modified Tmid, T90 or A42 or altered growth curve.

It is clear from the data as presented in the examples that one or more of the growth 35 characteristics as defined herein above, may be combined in one plant. Alternatively, depending on the chosen transformation event and/or depending on the promoter used, one

or more of these growth characteristics may be present or absent or more or less pronounced in the plant.

5 The methods of the present invention may also be used to confer stress tolerance to plants. In particular, a 2xC2H2 of the STZ type may be used to confer to a plant salt stress tolerance and/or drought stress tolerance. According to a specific embodiment, a tissue preferred promoter, such as a seed-preferred promoter" is used in these methods.

10 The present invention also relates to use of a nucleic acid sequence encoding a zinc finger protein and homologues, derivatives and active fragments thereof in modifying the growth characteristics of plants, preferably in increasing yield, further preferably increasing seed yield.

15 The present invention also relates to use of a nucleic acid sequence encoding a 2xC2H2 zinc finger protein and homologues, derivatives and active fragments thereof and to the 2xC2H2 zinc finger protein itself and to homologues, derivatives and active fragments thereof as a growth regulator. The sequences represented by SEQ ID NO 1, and portions thereof and SEQ ID NO 2, and homologues, derivatives and active fragments thereof are useful in modifying the growth characteristics of plants, as hereinbefore described. The sequences would therefore find use as growth regulators, such as herbicides or growth stimulators. The present invention also provides a composition comprising a protein represented by SEQ ID NO 2, or a homologue, derivative or active fragment thereof for the use as a growth regulator. A growth regulator is used herein as meaning a regulator that increased yield and is therefore also referred to as yield regulator.

20 In particular, the present invention provides a yield regulating composition comprising a nucleic acid encoding a 2xC2H2 protein, and/ or comprising a 2xC2H2 protein, and/or comprising a 25 construct as defined herein above. Such a yield regulating composition further comprises additives normally use in yield regulating compositions, such as a solvent or carrier.

30 Conversely, the sequences according to the present invention may also be interesting targets for agrochemical compounds, such as herbicides or growth stimulators. Accordingly, the present invention encompasses use of a nucleic acid encoding a 2xC2H2 protein, of a 2xC2H2 protein and/or of a construct as defined in any of claims 20 to 22 as target for an agrochemical, such as a herbicide or a growth stimulator.

35 The methods according to the present invention may also be practised by co-expression of a gene encoding a 2xC2H2 zinc finger protein in a plant with at least one other gene that cooperates with the gene encoding a 2xC2H2 zinc finger protein. Such a gene may be a gene encoding a target protein of the 2xC2H2 zinc finger protein. Co-expression may be effected by cloning the genes under the control of a plant expressible promoter in a plant expressible

vector and introducing the expression vector(s) into a plant cell using *Agrobacterium*-mediated plant transformation. Therefore, the methods according to the present invention may result in plants having modified growth characteristics, particularly increased yield, as described hereinbefore in combination with other economically advantageous traits, such as further yield-enhancing traits, tolerance to various stresses, traits modifying various architectural features and/or biochemical and/or physiological features.

5 Since the plants of the present invention have excellent growth characteristics and have high yield, they are suitable for the production of enzymes, pharmaceuticals or agrochemicals. Also, 10 there are suitable to produce food or feed products.

The invention clearly extends to enzymes, pharmaceuticals or agrochemicals as well as food or feed products isolated from these plants.

Further a nucleic acid encoding a 2xC2H2 protein, a 2xC2H2 protein and/or the constructs of 15 the present invention may be used breeding programs aiming at the development of plants with increased yield.

Particularly, the use of allelic variants as defined above in particular conventional breeding programmes, such as in marker-assisted breeding is also encompassed by the present invention; this may be in addition to their use in the methods according to the present 20 invention. Such breeding programmes sometimes require the introduction of allelic variations in the plants by mutagenic treatment of a plant. One suitable mutagenic method is EMS mutagenesis. Identification of allelic variants then takes place by, for example, PCR. This is followed by a selection step for selection of superior allelic variants of the sequence in question and which give rise to altered growth characteristics in a plant. Selection is typically carried out 25 by monitoring growth performance of plants containing different allelic variants of the sequence in question, for example, SEQ ID NO 1. Monitoring growth performance may be done in a greenhouse or in the field. Further optional steps include crossing plants in which the superior allelic variant was identified with another plant. This could be used, for example, to make a combination of interesting phenotypic features

30 According to another type of breeding programme a DNA marker is identified which may be genetically linked to a gene capable of modifying expression of a nucleic acid encoding a 2xC2H2 zinc finger protein in a plant, which gene may be a gene encoding the 2xC2H2 zinc finger protein itself or any other gene which may directly or indirectly influence expression of 35 the gene encoding a 2xC2H2 zinc finger protein and/or activity of the 2xC2H2 zinc finger protein itself. This DNA marker may then be used in breeding programs to select plants having altered growth characteristics.

The methods according to the present invention may also be practised by introducing into a plant at least a part of a (natural or artificial) chromosome (such as a Bacterial Artificial Chromosome (BAC)), which chromosome contains at least a gene encoding a 2xC2H2 zinc finger protein, optionally together with one or more related gene family members. Therefore, according to a further aspect of the present invention, there is provided a method for modifying growth characteristics of plants by expressing in a plant at least a part of a chromosome comprising at least a gene encoding a 2xC2H2 zinc finger protein.

10 The present invention will now be described with reference to the following figures in which:

15 **Fig. 1** is a map of an expression vector for the expression in plants of a 2xC2H2zinc finger protein under the control of a GOS2 promoter. CDS1536 is the internal code for the *Arabidopsis thaliana* salt tolerant zinc finger (STZ) protein cDNA. The zinc finger protein expression cassette has a GOS2 promoter and a double terminator sequence (T-zein and T-rbcS-deltaGA) located within the left border (LB repeat) and the right border (RB repeat) of the Ti plasmid. Cloned within these T-borders are also a screenable marker and a selectable marker, each under the control of a constitutive promoter (Prom), followed by a terminator sequence (poly a and t-NOS). Furthermore, this vector also contains an origin of replication (pBR322 (ori + bom)) for bacterial replication and a selectable marker (Sm/SpR) for bacterial selection.

20 **Fig. 2A** shows digital images from a T1 positive line transformed with an STZ zinc finger transgene under control of a GOS2 promoter and **Fig. 2B** shows digital images of corresponding nullizygotes plants.

25 **Fig. 3** lists sequences useful in the methods of the present invention. SEQ ID NO 1 is an STZ encoding nucleic acid isolated from *Arabidopsis thaliana*; the start and the stop codon are highlighted in bold. SEQ ID NO 2 is the STZ protein sequence encoded by SEQ ID NO 1. In the STZ protein the nuclear localization signal also called the KRS motif or B-box is annotated (bold, italics, underlined), as well as the L-box (bold, underlined), the EAR motif (bold, italics), and the two C2H2 zinc finger domains with QALGGH motif (bold and boxed). SEQ ID NO 10 to SEQ ID NO 25 provides the sequences of various orthologs of the *Arabidopsis thaliana* STZ protein from other plant species. SEQ ID NO 26 to SEQ ID NO 35 provides the sequences of various paralogs (from *Arabidopsis*) of the STZ protein. SEQ ID NO 36 to SEQ ID NO 50 provides the sequences of related 2xC2H2 genes and proteins useful in the methods of the present invention.

Fig. 4 is a photograph of T3 plants grown in a greenhouse (A) or in a field (B). The photograph shows yield increase (especially in aboveground biomass and plant height) in subsequent generations of STZ transformed plants.

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Fig. 5 shows the binary vector for expression in *Oryza sativa* of the *Arabidopsis thaliana* STZ gene (CDS1536) under the control of a seed preferred WSI18 promoter (PRO0151). This vector contains a T-DNA derived from the Ti Plasmid, limited by a left border (LB repeat, LB Ti C58) and a right border (RB repeat, RB Ti C58)).

10 The zinc finger protein expression cassette has a WSI18 (PRO0151) promoter and a double terminator sequence (T-zein and T-rbcS-deltaGA) located within the left border (LB repeat) and the right border (RB repeat) of the Ti plasmid. Cloned within these T-borders are also a screenable marker and a selectable marker, each under the control of a constitutive promoter (Prom), followed by a terminator sequence (poly a and t-NOS). Furthermore, this vector also
15 contains an origin of replication (pBR322 (ori + bom)) for bacterial replication and a selectable marker (Sm/SpR) for bacterial selection.

Examples

20 The present invention will now be described with reference to the following examples, which are by way of illustration alone.

DNA Manipulation

Unless otherwise stated, recombinant DNA techniques are performed according to standard protocols described in Sambrook (2001) Molecular Cloning: a laboratory manual, 3rd Edition
25 Cold Spring Harbor Laboratory Press, CSH, New York or in Volumes 1 and 2 of Ausubel *et al.* (1988), Current Protocols in Molecular Biology, Current Protocols. Standard materials and methods for plant molecular work are described in Plant Molecular Biology Labfase (1993) by R.D.D. Croy, published by BIOS Scientific Publications Ltd (UK) and Blackwell Scientific Publications (UK).

30

Example 1: Gene Cloning

A gene encoding an STZ protein was amplified by PCR from an *Arabidopsis thaliana* seedling cDNA library (Invitrogen, Paisley, UK). After reverse transcription of RNA extracted from seedlings, the cDNAs were cloned into pCMV Sport 6.0. Average insert size of the bank was
35 1.5 kb, and original number of clones was of 1.59×10^7 cfu. Original titer was determined to be 9.6×10^5 cfu/ml, after first amplification of 6×10^{11} cfu/ml. After plasmid extraction, 200 ng of template was used in a 50 μ l PCR mix. Sequences of the primers used for PCR amplification

were, including the *attB* sites for Gateway recombination (in bold) were PRM3204 (sense, start codon in italics) 5' **GGGGACAAGTTGTACAAAAAAGCAGGCTTCACAATGGCG** CTCGAGGCTC 3' (SEQ ID NO 3) and PRM3205 (reverse, complementary stop codon in italics) 5' **GGGGACCACTTGTACAAGAAAGCTGGTAATTCC7AAAGTTGAAGTTGA** 5 3' (SEQ ID NO 4).

PCR was performed using Hifi Taq DNA polymerase in standard conditions. The PCR fragment (CDS1536) was amplified and purified using standard methods. The first step of the Gateway procedure, the BP reaction, was then performed, during which the PCR fragment 10 was recombined *in vivo* with the pDONR plasmid to produce, according to Gateway terminology, an "entry clone", p3359. PDONR was purchased from Invitrogen, as part of the Gateway technology.

Example 2: Vector construction for rice transformation with pGOS2::AtSTZ

15 The entry clone p3359 was subsequently used in an LR reaction with p0640, a destination vector used for rice transformation. This vector contains as functional elements within the T-DNA borders a plant selectable marker and a Gateway cassette intended for LR *in vivo* recombination with the sequence of interest already cloned in the donor vector. Upstream of this Gateway cassette lies the rice GOS2 promoter for constitutive expression of the zinc finger 20 gene (De Pater et al., Plant J. 2 (6) 837-844, 1992). After the recombination step, the resulting expression vector with the expression cassette CD4398 (Figure 1) was transformed into *Agrobacterium* strain LBA4404 and subsequently into plants. Transformed rice plants were allowed to grow and then examined for various parameters as described in Example 3.

25 **Example 3: Evaluation of T0, T1 and T2 transgenic rice plants transformed with pGOS2::AtSTZ (CD4398)**

Approximately 15 to 20 independent T0 transformants were generated. The primary T0 transformants were transferred from tissue culture chambers to a greenhouse for growing and harvest of T1 seed. Six events of which the T1 progeny segregated 3:1 for presence/absence 30 of the transgene were retained. For each of these events, approximately 10 T1 seedlings containing the transgene (hetero- and homo-zygotes), and approximately 10 T1 seedlings lacking the transgene (nullizygotes), were selected by PCR. Based on the results of the T1 evaluation three events were chosen, for further characterisation in the T2 generation, one event being very positive for a number of parameters, a second event being positive for a 35 number of parameters, but less pronounced, and a third event being neutral. Seed batches from the positive plants (both hetero- and homozygotes) in T1, were screened by monitoring marker expression. For each chosen event, the heterozygote seed batches were then selected

for T2 evaluation. An equal number of positives and negatives within each seed batch were transplanted for evaluation in the greenhouse (i.e., for each event 40 plants were grown of which there were about 20 positives for the transgene and about 20 negative). Therefore, the total number for the three events amounted to 120 plants for evaluation in the T2 generation.

5

T1 and T2 plants were transferred to the greenhouse and evaluated for vegetative growth parameters and seed parameters, as described hereunder.

(I) Statistical analysis of phenotypic characteristics

10 A two factor ANOVA (analyses of variance) was used as statistical model for the overall evaluation of plant phenotypic characteristics. An F-test was carried out on all the parameters measured, for all the plants of all the events transformed with the gene of interest. The F-test was carried out to check for an effect of the gene over all the transformation events and to verify an overall effect of the gene or "global gene effect". Significant data, as determined by
15 the value of the f-test, indicates a "gene" effect, meaning that the phenotype observed is caused by more than the presence or position of the gene. In case of the F-test, the threshold for significance for a global gene effect is set at 5% probability level.

20 To check for an effect of the genes within an event, i.e., for a line-specific effect, a t-test was performed within each event using data sets from the transgenic plants and the corresponding null plants. "Null plants" or "Null segregants" are the plants treated in the same way as the transgenic plant, but from which the transgene has segregated. Null plants can also be described as homozygous negative transformant plants. The threshold for significance for the t-test is set at 10% probability level. Within one population of transformation events, some
25 events may be under or above this t-test threshold. This is based on the hypothesis that a gene might only have an effect in certain positions in the genome, and that the occurrence of this position-dependent effect is not uncommon. This kind of gene effect may also be referred to as a "line effect of a gene". The p- value is obtained by comparing the t-value to the t-distribution or alternatively, by comparing the F-value to the f-distribution. The p- value stand
30 for the probability that the null hypothesis (null hypothesis being "there is no effect of the transgene") is correct.

(II) Vegetative growth measurements

35 The selected plants were grown in a greenhouse. Each plant received a unique barcode label to link unambiguously the phenotyping data to the corresponding plant. The selected plants were grown on soil in 10 cm diameter pots under the following environmental settings: photoperiod= 11.5 h, daylight intensity= 30,000 lux or more, daytime temperature= 28°C or

higher, night time temperature= 22°C, relative humidity= 60-70%. Transgenic plants and the corresponding nullizygotes were grown side-by-side at random positions. From the stage of sowing until the stage of maturity (which is the stage where there is no more increase in biomass) the plants were passed weekly through a digital imaging cabinet (examples of 5 pictures are shown in Figures 2A and 2B). At each time point digital images (2048x1536 pixels, 16 million colours) were taken of each plant from at least 6 different angles. The parameters described below were derived in an automated way from the digital images using image analysis software.

10 **(a) Aboveground area**

Plant above ground area was determined by counting the total number of pixels from aboveground plant parts discriminated from the background. This value was averaged for the pictures taken on the same time point from the different angles and was converted to a physical surface value expressed in square mm by calibration. Experiments show that the 15 aboveground plant area measured this way correlates with the biomass of plant parts above ground.

Results of the maximum above ground area values of the lines selected for T2 evaluation are summarized in Table 1. The plants of the best performing line showed an increase in biomass 20 of 34 %, compared to the nullizygotes.

When an F-test was carried out on all the plants of all the T2 events it became clear that the transgenic plants show a significant increase in above ground area, in average an increase of approximately 18%. A significant increase in above ground biomass is also displayed by STZ transformed plants grown under field conditions (see figure 4).

25 *Table 1: Aboveground area of STZ transgenic T2 plants. Each row corresponds to one event, for which the average maximum aboveground area (expressed in mm²) was determined for the transgenics (TR) and the null plants (null). The difference in absolute values between the transgenic population and the nullizygotes of each event are presented (dif.) as well as the 30 percentage of difference between the two populations (% dif). P stands for the probability produced by the t-test for each event. The last row presents the average numbers calculated from all the events. Here the p-value is produced by the F-test.*

Total above ground Area Max (mm ²)					
Line	TR	null	dif	% dif	p-value
CD4396 L1	63947	47606	16341	34	0.0021
CD4396 L2	42509	41342	1167	3	0.8063
CD4396 L3	41116	33687	7429	22	0.1107
Overall	49178	41657	7522	18	0.0047

(b) Plant height measurements

Plant height was determined by the distance between the horizontal lines going through the

5 upper pot edge and the uppermost pixel corresponding to a plant part above ground. This value was averaged for the pictures taken on the same time point from the different angles and was converted, by calibration, to a physical distance expressed in mm. Experiments showed that plant height measured this way correlate with plant height measured manually with a ruler.

10 The increase in plant height was displayed very clearly in STZ transformed plants when measured at the end of the vegetative growth (see figure 4A). Also, this parameter, was displayed by STZ transformed plants when grown in the field (see figure 4B) at the time of harvest.

15 ***(c) Total area cycle time measurements***

Plants were imaged weekly along the complete cell cycle and the maximum total area of the plants was determined as mentioned above. Total Area Cycle Time is the time when a plant reaches 90% of its maximum total area. This parameter is an indication of the duration of the vegetative growth.

20 Only in some transgenic lines there was an effect on cycle time. These few lines showed a prolonged vegetative growth.

(III) Measurement of seed-related parameters

The mature primary panicles were harvested, bagged, barcode-labelled and then dried for

25 three days in the oven at 37°C. The panicles were then threshed and all the seeds collected. The filled husks were separated from the empty ones using an air-blowing device. After separation, both seed lots were then counted using a commercially available counting machine. The empty husks were discarded. The filled husks were weighed on an analytical balance and the cross-sectional area of the seeds was measured using digital imaging. This 30 procedure resulted in the set of seed-related parameters described below.

(a) Total number of filled seeds per plant

The number of filled seeds was determined by counting the number of filled husks that remained after the separation step.

5 Total numbers of filled seeds per plant are summarized in Table 2. The t-test shows that for two events, transgenic plants produce 106% and 130% more filled seeds than the nullizygotes.

10 Table 2: Number of filled seeds of STZ transgenic T2 plants. Each row corresponds to one event, for which the average number of filled seeds was determined for the transgenics (TR) and the null plants (null). The difference in absolute values between the transgenic population and the nullizygotes of each event are presented (dif.) as well as the percentage of difference between the two populations (% dif). P stands for the probability produced by the t-test for each event. The last row presents the average numbers calculated from all the events. Here 15 the p-value is produced by the F-test.

Number of filled seeds					
Line	TR	null	dif	% dif	p-value
CD4396 L1	387.9	188.7	199.19	106	<0.0001
CD4396 L2	163.8	156.5	7.22	5	0.8382
CD4396 L3	236.9	102.9	133.98	130	0.0004
Overall	264.9	159.7	105.25	66	<0.0001

(b) Total seed weight per plant

The total seed weight was measured by weighing all filled husks harvested from a plant.

20 The total seed weight values of STZ transformed plants are summarized in Table 3. STZ transgenic plants produce significantly more seed weight than the corresponding nullizygotes. The difference in seed weight of the transgenics may be as high as 138% or higher.

25 Table 3: Total seed weight per plant of STZ transgenic T2 plants. Each row corresponds to one event, for which the average total seed weight (in gram) was determined for the transgenics (TR) and the null plants (null). The difference in absolute values between the transgenic population and the nullizygotes of each event are presented (dif.) as well as the percentage of difference between the two populations (% dif). P stands for the probability produced by the t-test for each event. The last row presents the average numbers calculated 30 from all the events. Here the p-value is produced by the F-test.

Total weight of seeds					
Line	TR	null	dif	% dif	p-value
CD4396 L1	9.8	4.5	5.25	116	<0.0001
CD4396 L2	3.4	3.3	0.1	3	0.908
CD4396 L3	6.1	2.6	3.56	138	0.0001
Overall	6.5	3.7	2.75	74	<0.0001

(c) Harvest Index

The harvest index in the present invention is defined as the ratio between the total seed yield and the above ground area (mm²), multiplied by a factor 10⁶.

5

The harvest index values of the STZ-transgenic plants are summarized in Table 4. STZ transgenic plants have a significant increase in harvest index. The increase in harvest index of the transgenic plants may be as high as 66%, when compared to the corresponding nullizygotes.

10

Table 4: *Harvest index of STZ transgenic T2 plants. Each row corresponds to one event, for which the average harvest index was determined for the transgenics (TR) and the null plants (null). The difference in absolute values between the transgenic population and the nullizygotes of each event are presented (dif.) as well as the percentage of difference between the two populations (% dif). P stands for the probability produced by the t-test for each event. The last row presents the average numbers calculated from all the events. Here the p-value is produced by the F-test.*

Harvest Index					
Line	TR	null	dif	% dif	p-value
CD4396 L1	149.1	90	59.11	66	<0.0001
CD4396 L2	74	73.4	0.55	1	0.9574
CD4396 L3	121.3	75.9	45.32	60	<0.0001
Overall	114.8	82.6	32.16	39	<0.0001

(d) Thousand kernel weight (TKW) of plants

20 Thousand Kernel Weight (TKW) is a parameter extrapolated from the number of filled seeds counted, and their total weight.

The weight values of thousand kernels of STZ transgenic plants are presented in Table 5. STZ transgenic plants have increased thousand kernel weight. The increase of TKW of transgenic plants may be as high as 6% when compared to the corresponding nullizygotes.

Table 5: Thousand kernel weight of STZ transgenic T2 plants. Each row corresponds to one event, for which the average TKW was determined for the transgenics (TR) and the null plants (null). The difference in absolute values between the transgenic population and the nullizygotes of each event are presented (dif.) as well as the percentage of difference between the two populations (% dif). P stands for the probability produced by the t-test for each event. The last row presents the average numbers calculated from all the events. Here the p-value is produced by the F-test.

TKW					
Line	TR	null	dif	% dif	p-value
CD4396 L1	25.2	23.8	1.46	6	0.0128
CD4396 L2	20.6	20.7	-0.14	-1	0.7963
CD4396 L3	25.5	24.5	0.99	4	0.0812
Overall	23.7	23	0.71	3	0.0213

10 (e) **Total number of seeds**

The total number of seeds per plant was measured by counting the number of husks harvested from a plant.

15 The total numbers of seeds per plant are summarized in Table 6. STZ transformed plants have an increase in total number of seeds. The increase of total number of seeds may be as high as 68%, when compared to the corresponding nullizygotes.

20 Table 6: Total number of seeds of STZ transgenic T2 plants. Each row corresponds to one event, for which the average total number of seeds was determined for the transgenics (TR) and the null plants (null). The difference in absolute values between the transgenic population and the nullizygotes of each event are presented (dif.) as well as the percentage of difference between the two populations (% dif). P stands for the probability produced by the t-test for each event. The last row presents the average numbers calculated from all the events. Here the p-value is produced by the F-test.

Total number of seeds					
Line	TR	null	dif	% dif	p-value
CD4396 L1	483.5	367.4	116.03	32	0.0146
CD4396 L2	353.9	327.5	26.42	8	0.5473
CD4396 L3	383.6	228.2	155.48	68	0.0009
Overall	406	312.5	93.52	30	0.0002

Conclusion

It may be concluded that vegetative growth is increased in the STZ transgenic plants when compared to the control non-transgenic plants, as reflected by parameters such as above

5 ground area, where the increase is above 20 %. This effect may be attributed to the expression of the STZ gene in the transgenic plants. Additionally, in some transformation events, the length of the vegetative growth is altered in the STZ transgenic plants. For those transformation events in which this effect occurs, in average the vegetative growth was prolonged with about 4 to 6 days, under the conditions tested.

10 Furthermore, yield was increased in STZ transgenic plants. Several seed parameters reflect this yield increase. The total number of seeds harvested was at least 100% higher in the transgenics than in the control plants, for those events showing a differential. For these events, there was also an increase in the total number of seeds of the transgenics, which increase was higher than 30 %. Seed filling in those transgenics was greatly improved, 15 reaching differences above 100% in the number of filled seeds.

Seed of the transgenic plants were also heavier, and probably bigger, as suggested by the higher values obtained for the thousand kernel weigh. The TKW parameter is a very stable parameter in rice cultivars, such as nipponbare, and in the growth conditions here used. This means that this parameter is not easily influenced and makes it an important yield parameter.

20 Therefore a TKW increase of 6 % represents a significantly increase in yield. Harvest index, another important yield parameter, was increased in the transgenic plants with more than 50 %.

In summary, based on the evaluation of STZ transgenic plants in the T1, T2 and further 25 generations, it may be concluded that the presence of an STZ transgene, has a positive effect on the size of the plant and/or its organs, as well as a positive effect on the final yield harvested.

(III) Root growth measurements

Transgenic plants are grown next to their corresponding non-transgenic null segregant in transparent pots. In average, for each construct comprising a particular promoter-2xC2H2 30 combination, a minimum of 5 independent transformation events are evaluated for root growth, root development and root architecture. Typically, per transformation event, 10 transgenics are compared to 10 nullizygotes. Root pictures are taken weekly during plant growth. The pictures are processed and analyzed to extract the values for the root parameters as detailed below. Statistical analysis as described above are applied to these data.

a) Root Area

Total root area is calculated from the summed number of pixels of each root images. A positive linear correlation between root area and dry weight and root biomass of the root has been previously established by similar experiments. Therefore, root area is a good approximation for 5 root biomass.

b) Root Length

The total perimeter of the roots of a plant is calculated as the sum of the perimeter of all roots in the images. A linear correlation between this measurement and root length has been 10 previously established. Thus, root length is extrapolated from the total root perimeter.

c) Root Width

Average root width of a plant is expressed as the ratio between the Root Area and the Root Length.

15 STZ transgenic plants of the invention show a superior performance when compared to control plants. Transgenic plants are altered in one or more the root parameters detailed above. In particular the transgenic have increased root biomass, for example due to increased root dry weight or area, and/or increased root length and/or increased root width.

20 **Example 4: Leaf Blade Width Measurement.**
Leaves of STZ transgenic plants appeared bigger and wider when compared to the corresponding control non-transgenic plants. To quantify the increase in leaf width, leaf blade width (length of transversal axe) of the flag leaf was measured with a ruler at the widest point 25 of the leaf, which is approximately at half of the length, in plants that have reached the end of the vegetative growth phase. The results shown in the Table 7, indicate that the increase in the leaf blade width in at least the event here measured was around 15 % when compared to the corresponding nullizygote.

30 **Table 7: Leaf blade width of STZ transgenic T2 plants. The average leaf blade width was determined for the transgenics (TR) and the null plants (null) of the selected event. The difference in absolute values between the transgenic population and the nullizygotes of the event is presented (dif.) as well as the percentage of difference between the two populations (% dif). P stands for the probability produced by the t-test .**

35

Leaf blade width					
Line	TR	null	dif	% dif	p-value
CD4396 L1	1.56	1.35	0.21	15	0.098

Example 5: Vector construction for rice transformation with pWSI18::AtSTZ

Vector construction for transformation with the pWSI18 (PRO0151) - AtSTZ (CDS1536)

5 cassette was carried out essentially as in example 2. The entry clone p3359, described earlier, was subsequently used in an LR reaction with p05653, a destination vector used for rice transformation. This destination vector contains as functional elements within the T-DNA borders a plant selectable marker and a Gateway cassette intended for LR in vivo recombination with the sequence of interest already cloned in the donor vector. A WSI18 10 promoter for seed preferred expression (PRO0151) is located upstream of this Gateway cassette. After the recombination step, the resulting expression vector with the expression cassette CD4398 (Figure 5) was transformed into *Agrobacterium* strain LBA4404 and subsequently this vector was transformed to *Oryza sativa* plants. Transformed rice plants were allowed to grow and then examined for various parameters as described in example 3.

15

Example 6: Evaluation of T0 and T1 transgenic rice plants transformed with the seed preferred expression cassette pWSI18::AtSTZ (CD4398)

Preparations of calli and of the *Agrobacterium tumefaciens* strain containing the expression vector with the CD4398 expression cassette, were carried out as described in example 3, as 20 were the calli transformation and plant regeneration.

Approximately 15 to 20 independent T0 rice transformants were generated. The primary transformants were transferred from tissue culture chambers to a greenhouse for growing and harvest of T1 seed. Events, of which the T1 progeny segregated 3:1 for presence/absence of 25 the transgene, were retained. For each of these events, approximately 10 T1 seedlings containing the transgene (hetero- and homo-zygotes), and approximately 10 T1 seedlings lacking the transgene (nullizygotes), were selected by monitoring marker expression.

Transgenic plantlets were grown next to control nullizygotes, seeds were harvested and thousand kernel weight determined as previously described.

30

Transformed plants comprising the expression cassette CD8490 (seed preferred pWSI18:STZ), had a normal and healthy appearance and were harvested at the same time as the control plants. The seeds harvested from the transgenic plants had an increase in

thousand kernel weight when compared to the control plants. As shown in Table 8 increase in thousand kernel weight was above 10%.

Table 8: Thousand kernel weight of STZ transgenic T1 plants. The average 1 thousand kernel

5 weight was determined for the transgenics (TR) and the null plants (null) of the selected event. The difference in absolute values between the transgenic population and the nullizygotes of the event is presented (dif.) as well as the percentage of difference between the two populations (% dif). P stands for the probability produced by the t-test.

Thousand kernel weight					
Line	TR	null	dif	% dif	p-value
CD8490 L1	29.6	26.8	2.82	11	0.001

10

Example 7: Cloning, transformation and evaluation of other 2xC2H2 encoding genes.

In Table 9 an overview is given of constructs with STZ or other 2xC2H2 zinc finger proteins, under control of various promoters, which constructs are made for use in the methods of the present invention. The coding regions of the 2xC2H2 genes to be cloned (GOI, Gene of Interest) are amplified by PCR from cDNA, following the protocol as in Example 1. Specific primers for each 2xC2H2 gene were designed at the start and stop codons of the gene sequence as present in the public database under the accession number as indicated in Table 9. These cloned sequences are also herein incorporated under the SEQ ID NO number as mentioned in the table. Moreover, the isolated PCR fragments were also given a unique CDS number.

20 The PCR fragment with a 2xC2H2 gene is then cloned under the control of a particular promoter. Different combinations for different genes are made (see Table 9). Chimeric constructs are made and CD numbers represent bacterial strains carrying the chimeric construct. Corresponding transgenic plants are obtained by transforming the plants with the 25 chimeric constructs, following the protocols as mentioned herein before. Evaluation of the transgenic events reveals an increase in yield, and increase in leaf surface area and/or an increase in duration of vegetative growth in the transgenic plants when compared to the control non-transgenic plants.

30

CD-070-PCT

Table 9: examples of 2xC2H2 chimeric constructs useful for the methods of the present invention. *see Table 8

CDS	Accession number (cDNA on which primers were designed to amplify the CDS region)	Prot ACC number	SEQ ID NO	PRO0129*	PRO0170*	PRO0061_2*	PRO0123*	PRO0287*	PRO0111*
CDS1536 STZ Arabidopsis	X95573	CAA64820	1 + 2	CD4398	CD11371	CD11382	CD10960	CD10959	CD10313
CDS2200 Paralog Arabidopsis	AF022658 NM_120516	AAB80922.1At5g04340	28 + 29	CD11576			CD11413		CD11540
CDS2205 Paralog Arabidopsis	NM_123683	At5g43170	32 + 33	CD11325			CD11414		CD11387
CDS2775 Ortholog Oryza sativa	AF332876	AAK01713.1	36 + 37	CD09948					CD10311
CDS1677 Homolog Arabidopsis	AL132966 REGION: 116202..116729	CAB67667	38 + 39	CD06462			CD		CD
CDS3337 Homolog Sugarcane	CA279020		40	CD			CD		CD
CDS2416 Homolog Arabidopsis	AF254447	At3g57670	41 + 42	CD			CD		CD
CDS2377 Homolog Arabidopsis	AJ311810	CAC86167	43 + 44	CD			CD		CD
CDS Homolog Arabidopsis	AL355775 REGION: complement(7857..8451)	CAB80935	45 + 46	CD			CD		CD
CDS Homolog Arabidopsis	AL391143 REGION: complement(31730..32938)	CAC01747	47 + 48	CD			CD		CD
CDS3641 Homolog Arabidopsis	X98678	CAA67236	49 + 50	CD			CD		CD

Table 10: examples promoters used in combination with 2xC2H2 for the methods of the present invention.

Promoter	Preferred expression type	Origin species	Gene
PRO0151	Seeds (mainly embryo and aleurone). Strong expression.	Oryza sativa	WSI18
PRO0110	Root	Oryza sativa	RCc3
PRO0207	Green tissue. Moderate expression levels	Saccharum officinarum	Prp
PRO0123	Green tissue. Strong expression levels.	Oryza sativa	Protochlorophyllide reductase
PRO0090	Seed specific (mainly endosperm)	Oryza sativa	Prolamin RP6
PRO0170	Constitutive. Strong Expression.	Oryza sativa	High Mobility Group protein
PRO0218	Seeds (mainly embryo and aleurone)	Oryza sativa	oleosine 18kda
PRO0061_2	Young expanding tissues	Oryza sativa	beta-expansine EXPB9
PRO0129	Constitutive. High expression levels.	Oryza sativa	GOS2

5

Example 8: use of the invention in corn.

The methods of the invention described herein are also used in maize. To this aim, an STZ encoding gene, for example a maize or other STZ ortholog, is cloned under control of a promoter operable in maize, in a plant transformation vector suitable for Agrobacterium-mediated corn transformation. Methods to use for corn transformation have been described in literature (Ishida et al., Nat Biotechnol. 1996 Jun;14(6):745-50; Frame et al., Plant Physiol. 2002 May;129(1):13-22).

Transgenic plants made by these methods are grown in the greenhouse for T1 seed production. Inheritability and copy number of the transgene are checked by quantitative real-time PCR and Southern blot analysis and expression levels of the transgene are determined by reverse PCR and Northern analysis. Transgenic lines with single copy insertions of the transgene and with varying levels of transgene expression are selected for T2 seed production.

20

Progeny seeds are germinated and grown in the greenhouse in conditions well adapted for maize (16:8 photoperiod, 26-28°C daytime temperature and 22-24°C nighttime temperature)

as well under water-deficient, nitrogen-deficient, and excess NaCl conditions. Null segregants from the same parental line, as well as wild type plants of the same cultivar are used as controls. The progeny plants resulting from the selfing or the crosses are evaluated on different biomass and developmental parameters, including, plant height, stalk/stem thickness, stem size, number of leaves, total above ground area, leaf greenness, time to maturity, time to silking, flowering time, time to flower, ear number, ear length, row number, kernel number, kernel size, kernel oil content, grain maturity, harvesting time. The seeds of these lines are also checked on various parameters, such as grain size, total grain yield per plant, and grain quality (starch content, protein content and oil content).

5

Lines that are most significantly improved compared to corresponding control lines are selected for further field-testing and marker-assisted breeding, with the objective of transferring the field-validated transgenic traits into commercial germplasm. The testing of maize for growth and yield-related parameters in the field is conducted using well-established protocols.

10

15 The corn plants are particularly evaluated on yield parameters, such as for example, amount of plants per acre, amount of ears per plant, amount of rows per ear, amount of seeds per row and TKW. Subsequent improvements for introgressing specific loci (such as transgene containing loci) from one germplasm into another is also conducted using well-established protocols.

20

Claims

1. Method for increasing plant yield relative to corresponding wild type plants, comprising modifying expression in a plant of a nucleic acid sequence encoding a 2xC2H2 zinc finger protein and/or modifying in a plant level and/or activity of a 2xC2H2 zinc finger protein.
- 5 2. Method for increasing leaf surface area relative to corresponding wild type plants, comprising modifying expression in a plant of a nucleic acid sequence encoding a 2xC2H2 zinc finger protein and/or modifying in a plant level and/or activity of a 2xC2H2 zinc finger protein.
- 10 3. Method for prolonging vegetative growth phase of a plant relative to corresponding wild type plants, comprising modifying expression in a plant of a nucleic acid sequence encoding a 2xC2H2 zinc finger protein and/or modifying in a plant level and/or activity of a 2xC2H2 zinc finger protein.
- 15 4. Method according to any of claims 1 to 3, wherein said modifying expression, level and/or activity is effected by recombinant means and/or chemical means.
- 20 5. Method according to any of claims 1 to 4, wherein said 2xC2H2 zinc finger protein comprises a QALGGH motif.
6. Method according to any of claims 1 to 4, wherein said 2xC2H2 zinc finger protein comprises a NNM(W)QMH motif.
- 25 7. Method according to any of claims 1 to 6, wherein said 2xC2H2 zinc finger protein comprises an EAR motif.
8. Method according to any of claims 1 to 7, wherein said 2xC2H2 zinc finger protein further comprises a B-box.
- 30 9. Method according to any of claims 1 to 8, wherein said 2xC2H2 zinc finger protein further comprises an L-box.
- 35 10. Method according to any of claims 1 to 9, wherein said 2xC2H2 zinc finger protein is derived from a dicotyledonous plant, preferably from the family *Brassicaceae*, further preferably from *Arabidopsis thaliana*, more preferably the nucleic acid is as represented by

SEQ ID NO 2 or a homologue, derivative or active fragment thereof and/or wherein said nucleic acid is as represented by SED ID NO 1 or a portion thereof or sequences capable of hybridising therewith .

5 11. Method according to claim 10, wherein said homologue, derivative or active fragment has, in increasing order of preference, at least 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 52%, 54%, 56%, 58%, 60%, 62%, 64%, 66%, 68%, 70%, 72%, 74%, 76%, 78%, 80%, 82%, 84%, 86%, 88%, 90%, 92%, 94%, 96%, 98% sequence identity with the sequence of SEQ ID NO
10 2.

12. Method according to any of claims 1 to 11, wherein said plant is a monocot.

13. Method according to any of claims 1 to 12, wherein said modifying expression is effected
15 by introducing into a plant a nucleic acid capable of modifying expression of a gene encoding a 2xC2H2 zinc finger protein and/or capable of modifying level and/or activity of a 2xC2H2 zinc finger protein.

14. Method according to claim 13, wherein said nucleic acid capable of modifying expression is
20 a nucleic acid encoding a 2xC2H2 protein, such as a 2xC2H2 protein as defined in any of claims 5 to 11.

15. Method according to claims 13 or 14, wherein said nucleic acid introduced into a plant is an alternative splice variant of a nucleic acid as defined in claim 14.
25

16. Method according to claims 13 or 15, wherein said nucleic acid introduced into a plant is an allelic variant of a nucleic acid as defined in claim 14.

17. Method according to claims 13 or 16, wherein said nucleic acid introduced into a plant is
30 comprised on at least part of a chromosome.

18. Method according to any of claims 1 to 17, wherein said modifying expression comprises increased expression.

35 19. Method according to any of claims 1 to 18, wherein expression of said nucleic acid is driven by a plant promoter, preferably a constitutive promoter, such as a GOS2 promoter.

20. Method according to any of claims 1 to 18, wherein expression of said nucleic acid is driven by a plant promoter, preferably a tissue preferred promoter, such as seed-preferred promoter.

5 21. Method according to any of claims 1 to 20, wherein said increased yield comprises increased above ground biomass.

22. Method according to any of claim 1 to 20, wherein said increased yield comprises increased seed yield.

10 23. Method according to any of claim 1 to 20, wherein said increased yield comprises increased root yield.

24. Construct comprising:

15 (i) A nucleic acid capable of modifying expression of a nucleic acid encoding a 2xC2H2 zinc finger protein and/or capable of modifying level and/or activity of a 2xC2H2 zinc finger protein;
(ii) One or more plant control sequence capable of driving expression of the nucleic acid sequence of (i); and optionally
20 (iii) A transcription termination sequence.

25. Construct according to claim 24, wherein said nucleic acid of (i) is a nucleic acid as defined in any of claims 14 to 17.

25 26. Construct according to claim 24 or 25, wherein said control sequences of (ii) is at least a constitutive promoter, such as a GOS2 promoter.

27. Construct according to claim 24 or 25, wherein said control sequences of (ii) is at least a tissue preferred promoter, such as seed-preferred promoter.

30 28. Host cell comprising a construct according to any of claims 24 to 27.

29. Method for the production of a transgenic plant having increased yield, increased leaf surface area and/or prolonged vegetative growth, which method comprises

35 (i) introducing into a plant or plant cell a 2xC2H2 zinc finger nucleic acid;
(ii) Cultivating the plant or plant cell under conditions promoting plant growth.

30. Plant obtainable by a method according to any of claims 1 to 23 and 29, which plant has increased yield, modified leaf surface area and/or prolonged vegetative growth, relative to corresponding wild type plants.

5

31. Transgenic plant having increased yield, increased leaf surface area and/or prolonged vegetative growth, which transgenic plant has modified expression of a nucleic acid encoding a 2xC2H2 zinc finger protein and/or modified level and/or activity of a 2xC2H2 zinc finger protein, relative to corresponding wild type plants.

10

32. Plant part, preferably a harvestable part, a propagule or progeny of a plant as defined in claim 30 or 31, which progeny has modified expression of a nucleic acid encoding 2xC2H2 zinc finger protein and/or modified level and/or activity of a 2xC2H2 zinc finger protein, relative to corresponding wild type plants.

15

33. Plant or plant part according to any of claims 30 to 32, which plant is a monocotyledonous plant, preferably a cereal.

20

34. Plant or plant part according to any of claims 30 to 33 selected from rice, maize, wheat, barley, millet, oats, rye, sorghum, soybean, sunflower, canola, sugarcane, alfalfa, leguminosae (bean, pea), flax, lupinus, rapeseed, tobacco, tomato, potato, squash, papaya, poplar and cotton.

25

35. Use of a nucleic acid encoding a 2xC2H2 protein, of a 2xC2H2 protein and/or of a construct as defined in any of claims 24 to 27 to increase plant yield.

30

36. A yield regulating composition comprising a nucleic acid encoding a 2xC2H2 protein, and/or comprising a 2xC2H2 protein, and/or comprising a construct as defined in any one of claims 24 to 27.

35

37. Use of a nucleic acid encoding a 2xC2H2 protein, of a 2xC2H2 protein and/or of a construct as defined in any of claims 24 to 27 to increase leaf surface area.

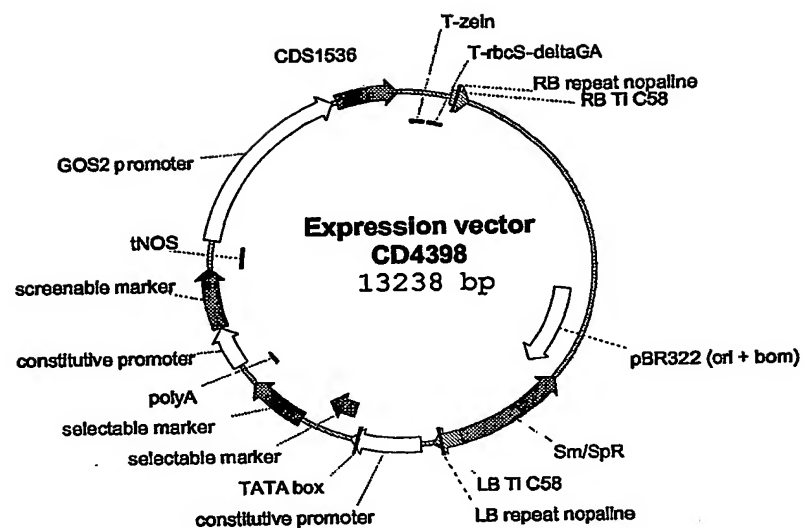
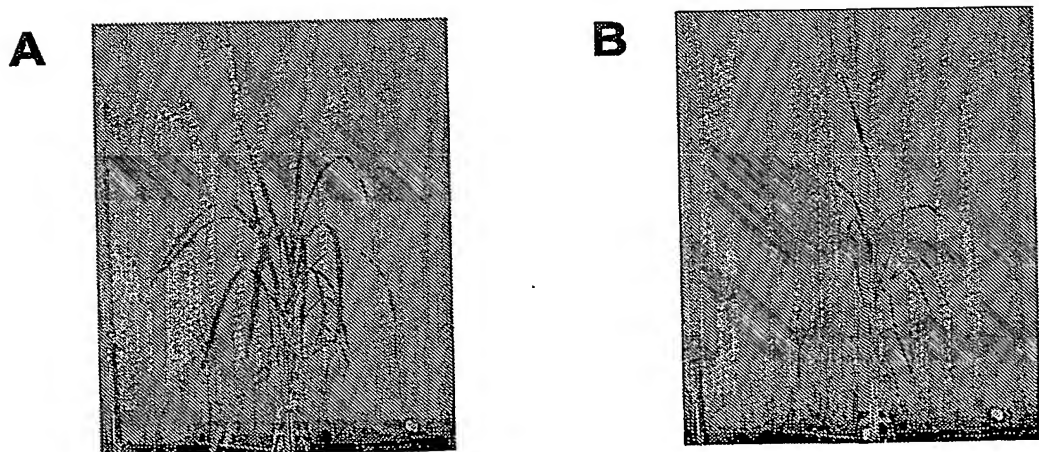
35

38. Use of a nucleic acid encoding a 2xC2H2 protein, of a 2xC2H2 protein and/or of a construct as defined in any of claims 24 to 27 to prolong vegetative growth.

39. Use of a nucleic acid encoding a 2xC2H2 protein, of a 2xC2H2 protein and/or of a construct as defined in any of claims 24 to 27 as target for an agrochemical.
40. Use of a nucleic acid encoding a 2xC2H2 protein, of a 2xC2H2 protein and/or of a construct as defined in any of claims 24 to 27 in a breeding program.
- 5 41. Use of a plant as defined in any of claims 30 to 34 to produce enzymes, pharmaceuticals or agrochemicals.
- 10 42. Use of a plant as defined in any one of claims 30 to 34 to produce food or feed products.

15

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**FIGURE 1****FIGURE 2**

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SEQ ID NO 1: *Arabidopsis thaliana* STZ cDNA (CDS1536)
 AATGGCGCTGAGGCTTACATCACCAAGATTAGCTTCCTCGATTCCTCTTGTGAG
 ATTCTCAGTCTTCCATGGAGTCGAGCACTGGACAAAGGTAAGCGATCTAAGAGATCAAGA
 TCCGATTCCACCACCAAAACCTCACTGAGGAAGAGTATCTAGCTTTGCCTCATGCTTCT
 CGCTCGCGACAACCGTCAAGCTCCTCCCTCCGGCGGTGGAGAAGTTGAGCTACAAGTGT
 GCGTCTCGCGACAAGACGTTCTTCTACCAAGCTCTCGGTGGTACAAGGCAAGCCACCGT
 AAGAACTTATCACAGACTCTCTCCGGCGGAGGAGATGATCATTAACCTCGTCGGCGACAAC
 CACATCCGCGTGAACACTGGAAGTGGGAAATCACACGTTGCACCATCTGTAACAAGTCTT
 TTCCCTCCGGTCAAGCTCTCGCGGACACAAGCGGTGCCACTACGAAGGAAACAACACATC
 AACACTAGTAGCGTGTCCAACCTCCGAAGGTGGGGTCCACTAGCCACGTTAGCAGTAGCCA
 CGTGGGGTTGACCTCAACATCCCTCGATCCCTGAATTCTCGATGGTCAACGGAGACGACG
 AAGTCATGAGCCCTATGCCGGCGAAGAAGCCTCGGTTGACTTCGGTCAAACCTCAACTT
TAAGGAAATT

SEQ ID NO 2: *Arabidopsis thaliana* STZ protein with annotation of the domains

B-BOX	L-BOX
MALEALTSPRLASPIPPLFEDSSVFHGVEHWTKG <u>KRSKRS</u> RSDFHHQNLTEEEYLA <u>FCL</u> MLL	
ARDNRQPPPPP <u>AVEKLSYK</u> CSVCDKT <u>FSSYQALGGH</u> KASER <u>RKNL</u> SQTLGGDDHSTSSATT	
TSAVTTGSGKSHV <u>CTICNKSFP</u> SG <u>QALGGH</u> KR <u>CEYEG</u> NNNINTSSVSNSEGAGSTSHVSSSH	
RGFD <u>LNIPPI</u> PEFSMVNGDDEVMS <u>PMPAKK</u> PR <u>FDPV</u> KLQL	

EAR motif

SEQ ID NO 3: PRM3204 (sense, start codon in italics)
 5' GGGGACAAGTTGTACAAAAAAAGCAGGCTTCACAATGGCGCTCGAGGCTC 3'

SEQ ID NO 4: PRM3205 (reverse, complementary stop codon in italics)
 5' GGGGACCACTTGTACAAGAAAGCTGGGTAATTCCTTAAAGTTGAAGTTGA 3'

BOXES

SEQ ID NO 5: QALGGH motif
QALGGH

SEQ ID NO 6: NNM motif
 NN (M/W) QMH

SEQ ID NO 7: EAR motif
 hDLNh (X) P

SEQ ID NO 8: B-Box
 KR (S) KRXR

SEQ ID NO 9: L-Box
 EXEXXAXCLXXL

FIGURE 3

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ORTHOLOGS OF STZ in OTHER PLANT SPECIES

SEQ ID NO 10: Dg_AF119050_1 *Datisca glomerata* zinc-finger protein 1 (zfpl) mRNA, complete cds

GGCACGAGGACAAATTCTCTCTATCCTCTGAATATCTTGGTTGTGAAC TGAGAAGCTA
 TTAGATGGCTCTAGAAGCGCTCAACTCTCCGACCACAGCTACGCCGGTGTTCACTACGACG
 ACCCCAGCTTGAATTACCTTGAGCCATGGACCAAGCGTAAGCGTTCCAAGCGTACCGCTTA
 GATAGCCCCATACCGAGGAAGAGTACCTGCTTCTGCCATCATGCTCGCTCGTGGCCGC
 GTTGCCTCTGCAAATCGACGGGATTCTCAGTCTCATTAGCATTAGCCTGAAGCAACGAC
 TTCGGCTACCAAAGTCAGTTATAAGTGCTCTGTGCGATAAGGCCCTTCTGCTTATCAGG
 CTTTGGGGGGACAAGGCCAGCCACAGAAAGCTCGCTGGCGCGAAGATCAATCGACTTCC
 TTTGCCACCACGAATTAGCCACCGTCACTACCACACAGCCTCCGGAGGTGGCAGGTC
 TCATGAGTGTCTATTGCCACAAATCGTCCCCACTGGCCAGGCCCTGGGTGGTCACAAGC
 GCTGCCACTACGAAGGCAGTATCGGCCGAATAGTATTACCACCAACAACAAATACCACCAAC
 AGCGGAAGCAACGGTGGCATGAGCATGACCTCCGAAGTAGGTTCCACACACAGTCAGCCA
 CAGTCACCGTGACTTCGATCTCAACATCCCGCCTGGCGAGTTCGGTGAATTCTTCA
 TATCCGGGGATGACGAGGTCGAGAGTCTCATCCGCCAAGAAACCCGTATATTGATGAAA
TAAAACATTCTCAAGATCACTGAACCAGGCTTAGTTCTTATAGGAGGAGATTAAAAAA
AGTAGTATCTCTTTCTTATCCGTAGGATAATTAATATATTCTGTACATAAAATTGTA
GTTCTTAACACACTCTGTTCACTTCTTGCTCAACTTGTATTGGTTATTCATT
ATGAAAATTCAATT

SEQ ID NO 11: Dg_AF119050_1 *Datisca glomerata*, STZ ortholog, protein

MALEALNSPTTATPVFHYDDPSLNYLEPWTKRKRSKRTRLDSPHTEEEYLAFCILMLARGV
 ASANRRDSQSSIQIPEATTSAKVSYKCSVCDKAFSSYQALGGHKASHRKLAGGEDQSTS
 ATTNSATVTTTASGGGRSHECSICHKSPTGQALGGHKRCHYEWSIGGNSIHHHNNTNS
 GSNGGMSMTSEVGSTHTVSHSHRDFDNI PALPEFRSNFFISGDDEVESPHPAKPRILMK

SEQ ID NO 12:Gm_T09602_U68763.1_GMU68763 *Glycine max* (soybean) probable zinc finger protein SCOF-1 mRNA, complete cds

AAAATTCTCACTCTCTCATCTCGAGATCATAGTATCATATTCAATATCATTCTACACC
 AAACACATGGCTTGGAGCTCTCAACTCACCAACAACAACCGCTCCATCTTCCCTTGA
 CGACCCAACATTCATGGCGAACGAAAACGTTCAAGCGTTCTCGCGACCATCCTCTG
 AAGAAGAGTACCTCGCCCTCTGCCCATCATGCTCGCTCGCCGGCACCACCGTCAAC
 AACCGCACGTCAGCCCTCGCCGCTACAGCCACAGCCACAGCCACACCAGATCCTCCAC
 CAAGCTCAGTTACAAATGCTCCGTTGCGACAAGAGCTTCCCTTACCAAGCGCTCGGTG
 GACACAAGGCCAGTCACCGGAAACTCGCCGGCCGCGAAGACCAACCCCCCAGCACCACC
 ACTTCCCTCCGCCGCCACAGCTCCGCTCCGGAGGTAAAGGCCATGAGTGTCCATTG
 CCACAAATCTCCCCACCGGACAGGCCCTGGCCGACACAAACGTTGTCACTACGAAGGTA
 ACGGTAACGGAAATAACAACACAGTAACAGCGTTGTCAACCGTCGCTCGGAAGCGTG
 TCCACCCACACTGTCAGTCACGGCCACCCACCGCAGCTCGATCTCAACATCCGGCTTCC
 GGATTTTCGACCAAGGTGGAGAAGACGAGGTTGAGAGGCCCTCACCTGTCAAGAAGC
 CTCGCCTTCGTCATTCCAAGATCGAAATCCCCAATTCAATGAACCTCGTTGAATT
 AGTTATTTTCGACTATATATTGGAGAATTGAGAGTTACTATAATTGATTTGTAC
 ATAGTACTTGGAGTTGTTGGACCGTACCGGACCCAGTTCTGGTTGAGGTTGTACTTT
 CACAACAGTGGCAGATTGCAATTCAATTATTGTTATTAAAAAA

FIGURE 3 (continued)

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SEQ ID NO 13: Gm_T09602 Glycine max (soybean) probable zinc finger protein SCO1-1, STZ ortholog, protein
 MALEALNSPTTAPSFPFDDPTIPWAKRKRSKRSRDHPSEEEYLALCLIMLARGGTTVNNR
 HVSPPPLQPQPQPTPDPSTKLSYKCSVCDKSFPSYQALGGHKASHRKLAGAAEDQPPSTTS
 SAAATSSASGGKAHECSICHKSFTPQALGGHKRCHYEGNGNGNNNSNSVVTASEGVGST
 HTVSHGHHRDFDLNIPAFPDFSTKVGDEVESPHPVMKKPRLFVIPKIEIPQFQ

SEQ ID NO 14: Ms_CAB77055_Y18788.1_MSY18788 Medicago sativa putative TFIIIA (or kruppel)-like zinc finger protein mRNA
 AATTCCGGCACGAGAAATAACCACTTCCTCTCAAAACCTCCTTGCCTTTGCTTCTACTT
 TCACTTGCCTAACGCTAACTAACTCTCGAGTGTCTCTTTCATCATATGGCTATGGA
 AGCACCTTAACCTACCCACCCTGCTACTCCTTCACACCCCTTGAGGAACCAAATCTGAGTT
 ATCTTGAACACCCGTGGACGAAAGGTAAACGATCAAAGCGTTCTCGCATGGATCAATCTCA
 TGCACTGAAGAAGAGTATCTGCTCTTGTCTCATCATGCTTGCTCGCAGCGGTAACAACAA
 CGACAAAAAGTCTGATTGGTGGCGACGCCGCTAACACCCTTAAACTCAGTCACAAATGCT
 CAGTCTGCAACAAAGCTTCTCATCTATCAAGCCCTAGGTGGACACAAAGCCAGTCACCGG
 AAAGCTGTTATGTCGCACCCACCGCTGAAGATCAGATCACCACCACTTCATCCGCCGTGAC
 TACCAAGCTCTGCTCCAACGGTAAGAACAAAGACTCATGAGTGTCCATCTGTACAAACCT
 TCCCTACTGGACAGGGCTTGGGAGGACACAAGCGTTGTCACTACGAAGGCAGCGTTGGGCC
 GGTGCCGGTGTGGAAGTAACGCTGTAACGCTCTGAAGGAGTTGGATTGTACACAGCCA
 CCACCGTGATTTGATCTAACCTCCGGCTTTCCGGACTTTCAAAGAAGTTTCTGTGG
 ATGACGAGGTTTAGTCCTTACCTGCTGCAAAGAACCCGTGTCTTTCAAGCTGGAAATT
 CCTTCTCATTACTGATCAATAATAGATCCAATTTTATTGTTATTATTATAATAATTATT
 CGCTTAGGGCATAGTTATTTCTTTCAATTATTCGGATCAATTGTTCTGTACA
 TACAAATTGGGATTGGTTAGAATTAGGACGGTTGTAGACAATGGAAATTCAATTCAATT
 ATTTAATTGTGT

SEQ ID NO 15: Ms_CAB77055_Medicago sativa putative TFIIIA (or kruppel)-like zinc finger protein, STZ ortholog, protein
 MAMEALNSPTTATPFTPFEENLSYLETPWTGKRSKRSRMDQSSCTEEYLALCLIMLARS
 GNNNDKSDSVATPLTTVKLSHKCSVNCNAFSSYQALGGHKASHRKAQMSATTAEDQITTS
 SAVTSSASNGKNTHECSICHKSFTPQALGGHKRCHYEGSVGAGAGAGSNAVTASEGVGL
 SHSHHRDFDLNLPAPDFSKKFFVDEVFSPLPAAKPCLFKLEIPSHY

SEQ ID NO 16: Nt_AAC06243_AF053077 Nicotiana tabacum osmotic stress-induced zinc-finger protein (zfp) mRNA, complete cds
 TTTTCCCTCGAATTGATAACTAAAGAGAATATTATGACTCTTGAAGCTTGAAAGTCACCTA
 CGGCGGCAACGCCGACTTACCAACCGCTATGAAGATGATGAAATTCTATAATTGGAT
 TCTGGGCTAAAGGAAACGATCAAACGGCCCCGTATTGATGCCAACCGACTGAAGAAGA
 GTATTTAGCCCTGTCTCATCATGCTCGCTCGCAGCGGAACCGGAACCGAGAACCGGTTAA
 CTGATGCTACTACTTCCAAACAACCTGCCGATAAAAAACCGCCGAGTTGCCGCCGGTTCAT
 AAGAAAGAGGTGGCAACAGAGCAAGCAGAGCAATCTTACAAGTGTAGCGTGTGACAAGGC
 TTTTCTTCTTATCAAGCACTCGGTGGGCATAAGCAAGTCACCGTAAACTACTACTG
 CTACCGCCGCTCTGATGATAACAATCCTCAACTCAACTTCCACTGGCGCCGTTAATATC
 TCTGCTCTTAATCCAATGGTCGTTCACACGTCTGTTCTATTGCCACAAGGCTTCTAC
 TGGCCAAGCTTGGGTGGCACAAGCGCCGCAACTATGAAGGCAAACTCGGTGGTAACAGCC
 GCGACTTAGGCGCGCGCGCGCGGTAGTGGAAAGCGTCTGACTACTTCAGACGGC

FIGURE 3 (continued)

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GGCGCGTCGACTCACCGCTACGTGACTTGACCTGAACATGCCTGCTTCGCCGGAATTGCA
 ACTGGGTCTGAGTATTGATTGTGGACGGAAAAGTCAACTGTTGCCGATGGTCCAAGAGGTGG
 AAAGTCCTATGCCTGCAAAGAAACCGCGTTATGTTCTGGTTTGAAACTCTTTAGG
 GGAATTGAATTGATTGTGTTAGCAAATTAGTAAATTGGTTCATGTGATTATTTAG
 GAAAAGGAATTATTGATTGTGTTACCGTTATTCTAGGGTGGTATTATGTACAGGGAGTG
 AATCATTCAATTGGTTTACACTTCTTAATTATATATTCTTTTACACATAAAAAAAA
 AAAAAAA

SEQ ID NO 17: *Nt_AAC06243_Nicotiana tabacum osmotic stress-induced zinc-finger protein, STZ ortholog, protein*
 MTLEALKSPTAATPTLPPRYEDDDEIHNLDSWAKGKRSKRPRIDAPPTEEYIALCLIMLAR
 SGTGTRTGLTDATTSQQPADKKTAELPPVHKKEVATEQAEQSYKCSVCDKAFFSYQALGGHK
 ASHRKTTTATAASDDNNPSTSTGAVNISALNPTGRSHVCSICHKAFTGQALGGHKRRH
 YEGKLGGSNSRDLGGGGGGHSGSVLTTSDGGASTHLDLNFNLMPASPELQLGLSIDCGRKS
 QLLPMVQEVESEPMKAKKPRLLFSLG

SEQ ID NO 18: *Os_AF332876 Oryza sativa zinc finger transcription factor ZF1 mRNA, complete cds*
 AATTCCGGCACGAGGCCACACAGCAACCAAGCCAGCTGCCACACTAGCTTGAGGCGAGCGAGCG
 AAGCTTAGCTAGCGGATAGAACAAAGTCGTCGATCTGCTTGCTGTTGTGAATTGCGGTGG
 AACGATGTCGAGCGCGTGTCCATGGAAGCGCTCCACGCCGCGGTGCTCAAGGAGGAGCAGC
 AGCAGCACGAGGTGGAGGAGGCGACGGTCGTGACGAGCAGCAGGCCACGAGCGGGAGGAG
 GGCAGCACCTGCCAGGGTGGCGAAGCGGAAGCGGGTCGCGCCAGCGATCGGAGGA
 GGAGAACCTCGCGCTCTGCCCTCATGCTCGCCCGCGCCAGGTGCGGAGTCAAGTGCTCCGCTGC
 CGCCTCCGCTCTCGGCTTCGGCGCCCCCGCCGGCAGGTGCGGAGTCAAGTGCTCCGCTGC
 GGCAAGTCCTCAGCTCTACCAAGCGCTCGCGGCCACAAGACGAGCCACCGGGTCAAGCT
 GCGACTCCGCCCGCAGCTCCGTCTGGCTCCGCCCGTGCCTGCTGCCCTCCG
 CCGAGGACCGCAGCCACGTATCCACCGCCGCTCCTCCGACGGCATGACCAACAGA
 GTCCACAGGTGTTCCATCTGCCAGAAGGAGTCCCCACCGGGCAGCGCTCGCGGGCACAA
 GAGGAAGCACTACGACGGTGGCGTAGGCGCCGGCGCATCTCAACCGAGCTCTGG
 CCACGGTGGCGCCGAGTCGAGGTGGGAAGCTCCGCAACGCCAGTCCGCCACCCGGCG
 TTGACCTCAACCTCCGGCGTGCCTGGAGTTCTGTGTCGGCGCGTCTCAAGGGCAAGAA
 GATGTGGGACGAGGAGGAGGAGGTCCAGAGCCCCCTGCCCTCAAGAAGCCCCGGCTCTCA
 CCGGTAAATTCAAGCAGCTGCACGGATCCGATCCGTAGAGTTTGTCTAGGGAGTGAATT
 CAGTCGAAACACACTATTCTGTTGATTCTGTTGTGCCGTATTGTTAATTGTTCTGCTT
 TTGTACAGAGCAAGCGAGTGATAACATAGCCATACATACAGTCATACAGATAAGGTCTAGCT
 CTTCTGGTCTTGTAAACACTGGAACACTGTACCTGTATCTTACACTTTGTTCTTGACA
 GTCATATATTGTAGACCAAAAAAAA

SEQ ID NO 19: *Os_AF332876 Oryza sativa seedling zinc finger transcription factor ZF1, STZ ortholog, protein*
 MSSASSMEALHAAVLKEEQQQHEVEEATVVTSSATSGEEGGHLPQGWAKRKRSSRRQRSEEE
 NLALCLIMLARGGHHRVQAPPPLSASAPPAGAEFKCSVCGKSFSSYQALGGHKTRVKLPTPPAAPVLA
 PAPVAALLPSAEDREPATSSTAASSDGMTNRVHRCSCICQKEFPTGQALGGHKR
 KHYDGGVGAGAGASSTELLATVAAESEVGSSGNGQSATRAFDLNLPAVPEFVWRPCS
 KGKMWDEEEEVQSPLAFKKPRLLTA

FIGURE 3 (continued)

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SEQ ID NO 20: Ph_BAA05079_D26086.1 [Petunia x hybrida].
PETZFP4 zinc-finger protein gene

TTCACTCACCAAAACAACCTCTCTACCTCTTACTTGACATTCAAATTCTTCATTACTA
CTTATCTCTACTAAATCTGATTGATTTAGTAAATCAAACAAGAGAATCTTTCAGTAATA
CAAACAAAGAAAATTTCTCTATACATTGATTGAGTTAGTAAGGCAAACAAGAAAATATC
ATGGCACTTGAAGCATTGAATTCTCCAACTACAAACAACACCACCATCATTCCAATTGAGAA
CAACGGGCTTAAGTACCTTGAGAGTTGGACAAAAGGTAAAAGATCAAAAGGCAACGCAGCA
TGGAACGACAGTGTACTGAAGAAGAGTATTGACTTTGCTTATCATGCTAGCACGTAGC
GATGGTTCTGTTAATAACTCACGGTCTTACCAACCACCACTACCACCATCAGTTCCAGT
AACGTCGCAAATAAACGCGACGTTATGGAACAGAAGAATTGTCAGTGTCCGTTGTG
GTAAAGGGTTGGGTCTTATCAAGCTTAGGTGGACATAAGCAAGTCACCGGAAACTGTC
AGCATGGGAGGAGATGAACAATCTACTACTTCCACTACTAACGTAACGGGAACTAGTTC
CGCTAACGTTAACGGTAACCGGAAGAACTCACGAATGTTCAATTGTCACAAGTGCTTCTA
CTGGACAAGCTTAGGTGGTCATAAAAGGTGCCACTATGACGGTGTACGGTAACGGTAAC
GGAAGTGTAAAGTGTGGGTGACGTATCTGAAGGTGTGGGTCCACTATTAGTCATCACCG
TGACTTGAATTGAAATATCCCGCGTTGCCGGAGTTGGCCGGATTGGTCCGGGAGG
ATGAGGTGGAGAGTCCTCATCCAGCAAAGAAGTCAGGCTATCTTCCACCTAAACTGAA
TTATTCAAAGGATTATAGAGGGAATTGATTGTTACAGGAAGATTATTAGGATTACGA
ATTTTTGTTGACTAGTTATGTAATAT

SEQ ID NO 21: Ph_BAA05079 [Petunia x hybrida] zinc-finger protein, STZ otholog, protein
MALEALNSPTTTTPPSFQFENNLKYLESWTKGKRSKRQRSMERQCTEEYLALCLIMLARS
DGSVNNRSRSLPPPLPPSVPVTSQINATLLEQKNLYKCSVCGKGFGSYQALGGHKASHRKL
SMGGDEQSTTSTTNVTGTSANVNGNGRTHECSICHKCFPTGQALGGHKRHYDGGNGNG
GSVSVGVTSSEGVGSTISHRDFDLNIPALPEFWPGFSGEDEVESPHPAKKSRLSLPPKLE
LFKGL

SEQ ID NO 22: Ta_BAA03901 Triticum aestivum gene for zinc-finger protein WZF1, complete cds
ATGTCGTCGCCATGGAAAGCGCTCCACGCCCTGATCCGGAGCAGCACAGCTGGACGT
TGAGGGGGCTGCCGCTGTCAGCAGGCCACCAGCGCGAGGGAGAGCGGCCACGTGCTGCAGG
GGTGGGCCAAGAGGAAGCGATCGCGCCGCCAGCGCTCCGAGGAGGAGAACCTCGCGCTCTGC
CTCCTCATGCTCTCGCGCCGGCAAGCAGCGTGTTCAGCGCCGCAGCCGGAGTCGTTCGC
TGCGCCGGTGCCGAGTTCAAGTGCTCCGTCTGCCGAAGTCCTTCAGCTCCTACCAGG
CGCTCGGAGGCCACAAGACGCCACCGGGTGAAGCAGCGTCTCCTCCCTGTATGCCGCT
GCTGCCCACTCGTGGCCCTCCCGGCCGTCGCCGCCATCCTGCCGCGCCAGCCGCCAC
GTCGTCCACCGCCCGTCCCTCGACGGCGCGACCAACAGAGTCACAGGTGCTCCATCTGC
AAAAGGAGTTCCGACTGGGCAGGCCGCTGGCCGGCACAAGAGGAAGCACTACGACGGAGGC
GTGGGCAGGCCGCGCCCTCGACCGAGCTCTGGCCGCCGCCGGCTTCGACCTGAACATTCCGGCGTGC
GAGCACCGGCAACGGGAGCTCGCCGCCGGCGAGTCGACCTGAACATTCCGGCGTGC
AGTCGTGTGGAGGCCGTGCACCAAGGGCAAGATGATGTGGGAGGACGATGAGGAGGTGCAG
AGCCCCCTCGCCTCAAGAAGCCTCGGCTTCTCACCGCTTGA

FIGURE 3 (continued)

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SEQ ID NO 23: Ta_BAA03901_WZF1 *Triticum aestivum*, STZ ortholog, protein
 MSSSAMEALHALIPEQHQLDVEAAAASSATSGEESGHVLQGWAKRKRSSRQRSEEEENLALC
 LLMLSRGGKQRVQAPQFESFAAVPAEFKCSVCGKSFSSYQALGGHKTSRVKQPSPPSDAA
 AAPLVALPAVAAILPSAEPATSSTAASSDGATNRVHRCSTCQKEPTGQALGGHKRKYDGG
 VGAAASSTELAAAEESEVGSTGNSSAARAFDLNIPAVPEFVWRPCAKGKMMWEDDEEVQ
 SPLAFKKPRLLTA

SEQ ID NO 24: Ca AF539746 *Capsicum annum* zinc finger protein mRNA, complete cds
 AAAATCTCGCTACTTACATCTAGAATAGTCACTAGAACAGTAACCTTATACAA
 CGGATATCGAT ATGGCACTTGAAGCTTGAATTCTCAACTGGTACACCAACTCCGCCACCG
 TTTCAATTGAGAGCGACGGCCAACAGCTTCGATATATCGAAAACGGAGGAAGGGAAAGAG
 ATCTAAAAGGTACCGCAGCATGGAGCACCGCCACTGAGGAAGAATACTTAGCGCTTGTT
 TGATCATGCTTGCACGTAGCGGTGGCTCGTTAACATCAACGATCTTACCCACCGCCGGCT
 CCGGTGATGAAACTGCACGCCGTGTCATCATGGCGGGAGGAGGAAGGAGAAGAT
 GGTGTATAAGTGTGGTTGTGGTAAGGGATTGGGTCTTATCAAGCTTGTGGACACA
 AAGCTAGTCACCGGAAACTCGTACCCGGCGAGATGATCAGTCAACTACCTCCACAACCACT
 AACGCAACCGGAAACAACACCTCCGTTAACGGCAACGGCAACAGAAGTGGAGGACTCACGA
 GTGTTGATTTGTCACAAGTGTGTTTCCACTGGACAAGCTTGTGGACACAAAAGGTGTC
 ACTACGACGGCGGTATCGGTAAACGGAAACGCTAACAGTGGCGTTAGTGCTAGCGTTGGAGTG
 ACGTCATCGGAGGGTGTGGGTCCACAGTCAGTCACCGGGATTCGACTTGAACATTCCGGC
 GTTGGCGGAATTCTGGCTGGGATTGGTCTCGCGAAGATGAGGTTGGAGAGTCCACATCCGG
 CGAAGAAATCGCGGTATGTTGCCTCCAAAATATGAATTATTCACACAT TAATGGGAATT
 GATTGTAGGATTACTATTTGGTAGACAAAATTATACTATGTAAGTTAATTTCATTG
 TGGTGGGAGCAAAATTAAATTGTCTATAGACCTAGCTAGTTACTAATAGCAAAA
 TTCAATTGATTGATTAAAAAAA

SEQ ID NO 25: Ca AF539746-*capsicum annum*, STZ ortholog, protein
 MALEALNSPTGTPPPFQFESDGQQLRYIENWRKGKRSRSRSMEHQPTEEYLALCLIML
 ARSGGSVNHQRSLPPPAPVMKLHAPSSSSAAEEEKEKMYKCSVCGKFGSYQALGGHKASH
 RKLVPGDDQSTTSTTNATGTTSVNGNGNRSGRTHECSICHKCFPTGQALGGHKRHYDG
 GIGNGNANSVGSASVGVTSEVGSTVSHRDFDLNIPALPEFWLGFSGSEDEVESPHPAKKS
 RLCLPPKYELFQH

PARALOGS OF STZ IN ARABIDOPSIS THALIANA

SEQ ID NO 26: gi_18402298_ref_NM_112848.1 mRNA
 ACTTCACTCTCTAATTCCCTCTCTATCTCTACCATATTGGGATTAAAAACTCTCAAC
 TTTTCTCTCAAATTCTGATCCTTGATCCAACAGTTAGAAGAAGATTCACTGATCATGGC
 CCTCGAAGCGATGAACACTCCAACCTCTTCTTCAACAGAATCGAAACGAAAGAAGATTGA
 TGAACGACGCCGTTTCATTGAGCGTGGCTAAACGCAAACGCTCCAAACGTCAGCGTTCT
 CACAGCCCTCTCGTCTTCTCACCGCCTCGATCTGACCCAAATCCAGAATCAAGA
 TCTTACGGAAGAAGAGTATCTCGCTTTGTCTCCTCATGCTCGTAAAGATCAACCGTCGC
 AAACGCGATTTCATCAACAGTCGCAATCGTTAACGCCGCCAGAATCAAAGAACCTCCG
 TACAAGTGTAAACGTCGTGAAAAAGCGTTCTTCTATCAGGCTTGTAGGCGGTACAAAGC
 AAGTCACCGAATCAAACCAACCGTAATCTCAACAAACGCCGATGATTCAACAGCTCCGA

FIGURE 3 (continued)

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CCATCTCCATCGCGCCGGAGAAAAACATCCGATTGCTGCCTCCGGAAAGATCCACGAGTGT
 TCAATCTGTCTAAAGTGTTCGGACGGGTCAAGCTTACGGCGGTACAAACGTTGTCACCA
 CGAAGGCAACCTCGCGCGAGGAGGAGGAGGAAAGCAATCAATCAGTCACAGTGGAAAGCG
 TGTCGAGCACGGTATCGGAAGAAAGGAGCCACCGTGGATTCATCGATCTAACCTACCGCG
 TTACCTGAACTCAGCCTTCATCACAATCCAATCGTCGACGAAGAGATCTGAGTCCGTTGAC
 CGGTAAAAAACCGCTTTGTGACCGATCACGACCAAGTCATCAAGAAAGAAGATTATCTT
 TAAAAATCTAATACTCGACTATTAAATTCTTGTGATTTTCGTTACAACCATAGTTCA
 TTTTCATTTTTAGTTACAAATTAAATTGTTGATTGAATATTGGTATATTG
 TTAGGGGTTGATAAC

SEQ ID NO 27: Translation of gi_18402298_ref_NM_112848.1
 MALEAMNTPTSSFTRIETKEDLMNDAVFIEPWLRKRSKRQSHSPSSSSSPRSRPKSQN
 QDLTEEEYLALCLLMLAKDQPSQTRFHQQSOSLTPPPESKNLPYKCNVCEKAFPSYQALGGH
 KASHRIKPPVTISTTADDSTAPTTISIVAGEKHPIAASGKIHECSICHKVFPTGQALGGHKRC
 HYEGNLGGGGGGGSKSISHSGSVSSTVSEERSHRGFIDLNLPALPELSLHHNPIVDEEILSP
 LTGKKPLLLTDHDQVIKKEDLSLKI

SEQ ID NO 28: gi_30680473_ref_NM_120516.3_mRNA
 AAATCAAATCTTCATTTACAATTATCTTCTCAATTAGAACCTTAGTAGCTAGCTTT
 CAAGATAATGGCACTTGAAACTCTTACTTCTCCAAGATTATCTCTCCGATGCCACTCTGT
 TTCAAGATTCACTAGGGTTTCATGGAAGCAAAGGCAAACGATCTAACGATCAAGATCT
 GAATTGACCGTCAGAGTCTCACGGAGGATGAATATATCGCTTATGTCATGCTTCTGC
 TCGCGACGGAGATAGAAACCGTGACCTTGACCTGCCTTCTTCGCTTACACCTCTGC
 TTCCCTCTTCCACTCCGATCTACAAGTGTAGCGTCTGTGACAAGCGTTTCGTTAC
 CAGGCTCTGGTGACACAAGGCAAGTCACCGAAAAGCTTTCGTTACTCAATCTGCCGG
 AGGAGATGAGCTGTCGACATCGTCGGCGATAACCACGTCGGTATATCCGGTGGCGGGGAG
 GAAGTGTGAAGTTCGACGTTGCTCTATCTGTCATAAAATCGTCCGACCGGTCAAGCTCTC
 GGCACAAACGGTGCCACTACGAAGGAAAGAACGGAGGCGGTGTGAGTAGTAGCGTGTG
 GAATTCTGAAGATGTGGGTCTACAAGCCACGTCAGCAGTGGCCACCGTGGTATGACCTCA
 ACATACCGCCGATACCGGAATTCTCGATGGTCAACGGAGACGAAGAGGTGATGAGTCCTATG
 CCGCGAAGAAACTCCGGTTGACTTCCGGAGAAACCTAAACATAAACCTAGGAAACT
 TTACAGAATTCTTATAGGAAATTGTTACTGTATATAACAAATATCGATTGATGTTCTGAAAAGATAT
 GTTCTCTTCACTGAAAATTATGATTCTTGTATAATTGATGTTCTGAAAAGATAT
 AACTTTTATTGTTCACACGTATCAAAATTGCTTGGATACATCA

SEQ ID NO 29: Translation of gi_30680473_ref_NM_120516.3
 MALETLTSPRLSSPMPTLFQDSALGFHGSKGKRSRSEFDRQSLTEDEYIALCLMILLARD
 GDRNRDLDLPSSSSSPPLPPLPTPIYKCSVCDKAFSSYQALGGHKASHRKSFSLTQSAGGD
 ELSTSSAITTSGISGGGGGSVKSHVCSICHKSFATGQALGGHKRCHYEGKNGGGVSSSVNS
 EDVGSTSHVSSGHRGFIDLNIPIPEFSMVNGDEEVMSPPMANKLRFDFPEKP

FIGURE 3 (continued)

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SEQ ID NO 30: gi_30693252_ref_NM_114853.2_mRNA

ATGGCTCTCGACACTCTCAATTCTCCCACCTCCACCACCAACCACCGCTCCTCCTCCTT
 CCTCCGTTGCCTCGACGAAACCGAGCCGAAAACCTCGAATCATGGACCAAAGAAAACGTA
 CAAAACGTCACCGTATAGATCAACCAAACCTCCTCCTCTGAAGAAGAGTATCTCGCTTT
 TGCCTCCTTATGCTCGCTGGCTCCTCCGATCATCACTCTCCACCGTCGGATCATCACTC
 TCTTCTCCACTGTCCGATCATCAGAAAGATTACAAGTGTCCGTCTGTGGCAAATCTTCC
 CGTCTTACCAAGCGTTAGGGACACAAAACAAGTCACCGGAAACCGGTTAGTGTGATGTT
 AATAATAGTAACGGAACCGTACTAATAACGGAATATTAGTAACGGTTAGTTGGTCAAAG
 TGGGAAGACTCATAACTGCTCTATATGTTAAAGTGTTCCTCTGGTCAAGCATTGGGTG
 GTCACAAACGTTGTCACTATGATGGGGTAACGGTAACAGTAACGGTACAATAGCCACAAG
 TTTGACCTAAATTACCGGCTGATCAAGTTAGTGTGAGACAATTGGAAAAGTCAACTCTC
 CGGTGAAGAAACAAAGTCGGTGTGTGATTATTATTACCGATCGGGATTAGCTAG
 TGGTTGATCATTAGCTGAGTCTGTAATGAAAATGAT

SEQ ID NO 31: Translation of gi_30693252_ref_NM_114853.2

MALDTLNSPTTTTAPPPFLRCLDETEPENLESWTKRKRTKRHRIDQPNPPSEEEYLAL
 CLLMLARGSSDHSSPPSDHHSPLSLSDHQDYKCSVCGKSFPSYQALGGHKTSRKPVSVDV
 NNSNGTVTNNGNISNGLVGQSGKTHNCSCIFKSFPSQALGGHKRHYDGGNGNSNGDNSHK
 FDLNLPADQVSDETIGKSQLSGEETKSVL

SEQ ID NO 32: gi_30694224_ref_NM_123683.2_mRNA

AAATTTCTATAGCAATGGCGCTTGAAGCTCTTAATTACCAAGATTGGTCGAGGATCCCTT
 AAGATTCAATGGCGTTGAGCAGTGGACCAAATGTAAGAAACGATCCAACGTTGAGATCTG
 ATCTTCATCATAACCACCGTCTCACTGAGGAAGAGTATCTAGCTTCTGTCTCATGCTCTT
 GCTCGGGATGGCGGCATCTGACTCTGTGACGGTTGCAGGAGAACGCCGAGTTATAAGTGTGG
 CGTTTGTACAAGACGTTTCGTCTTACCAAGCTCTCGGCGGTCTAAAGCGAGCCACCGGA
 GCTTATACGGTGGAGAGAATGATAAAATCGACACCATCCACCGCCGTGAAATCTCACGTT
 TGTTCGGTTGCGGGAAATCTTCGCCACCGGTCAGCTCTCGGCGGCCACAAGCGGTGCCA
 CTACGATGGTGGCGTTGAACTCGGAAGGTGTGGGTCTACTAGCCACGTCAGCAGTAGTA
 GCCACCGTGGATTGACCTTAATATTACCGGTGCAGGGATTTCGCCGGACGACGAAGTG
 ATGAGTCCGATGGCAGTAAGAAGCCTCGCTGAAGTAAGTCTTGTGAAGACCTGGAAGT
 TTATCAAATGAAATATCAAATTCAATTCAAGGAACAGTTGTGATTCTATTACCAAT
 ACACAATACGATTCAATTCC

SEQ ID NO 33: Translation of gi_30694224_ref_NM_123683.2

MALEALNSPRLVEDPLRFNGVEQWTKCKKRSKRSRSDLHHNHRLTEEEYLAFCLMLLARDGG
 DLDSTVVAEKPSYKCGVCYKTFSSYQALGGHKASHRSLYGGENDKSTPSTAVKSHVCVCG
 KSFATGQALGGHKRHYDGGVSNSEVGSTSHVSSSHRGFDLN1IPVQGFSPDDEVMSPM
 TKKPRLK

FIGURE 3 (continued)

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SEQ ID NO 34: gi_30698307_ref_NM_126145.2_mRNA

CACACTTCACTCTTCTTCATCTTCTTAAATAGCTCGAAATCACATCTCACAGAAAT
 TAAATCTATGGCTCTCGAGACTCTCAATTCTCAACAGCTACCACCAACCGCTCGGCCTCTT
 CTCCGGTATCGAAGAAATGGAGCCTGAGAATCTCGAGCAATGGCTAAAAGAAAACGAAC
 AAAACGTCAACGTTTGATCACGGTCATCAGAACTCAAGAAACGAACAAGAACCTCCTCTG
 AAGAAGAGTATCTCGCTTTGTCTCCTCATGCTCGTCGTGGCTCCGCGTACAATCTCCT
 CCTCTTCTCCTCACCGTCACGTGCGTCACCGTCCGATCACCGAGATTACAAGTGTACGGT
 CTGTGGGAAGTCCTTTCGTACATACCAAGCCTAGGTGGACACAAGACGAGTCACCGGAAAC
 CGACGAACACTAGTATCACTCCGTAACCAAGAACTGTCTAATAACAGTCACAGTAACAGC
 GGTTCGGTTATTAAACGTACCGTAACACTGGTAACGGTGTAGTCAAAGCGGAAAGAT
 TCACACTGCTCAATCTGTTCAAGTCGTTGCGTCTGGTCAGCCTAGGTGGACACAAAC
 GGTGTCACTATGACGGTGGCAACAACGGTAACGGTAACGGAAAGTAGCAGCAACAGCGTAGAA
 CTCGTCGCTGGTAGTGACGTCAGCGATGTTGATAATGAGAGATGGTCCGAAGAAAGTGCAT
 CGGTGGCCACCGTGGATTGACCTAAACTTACCGCTGATCAAGTCCTCAGTGAACGACTTCTT
 AACGTTGACTGAGTTGAGGAAAAGTCAACTATCAAGCGAAGAAAGGGTTAGTGGACGGTG
 AAGATTAACGGTCGTTCTTCCAGTGCTCGGTTGAGCTTGACTGGGTCTGTAATGAAA
 ATGATTGGAGTGGACTTGGCATTATTATTATTATTAAAAGAAATGTTAATTGTTGTT
 GGATTGTTATAGATAGAGGAAACAATTGGGATACACAAATTTTTTTTACAAGA
 AAATAATAATGCAGAGATGGATGATTGGATCGTACACGTATTATATAGTGGACCATTCTGT
 AATCGTGAATTATTATTGTTAGAAATTAAATTTCGT

SEQ ID NO 35: Translation of gi_30698307_ref_NM_126145.2
 MALETLSPTATTTARPLLRYREEMEPENLEQWAKRKRTKRQRFDHGHQNQETNKNLPSEEE
 YLALCLLMLARGSAVQSPPLPLPSRASPDSHDYKCTVCGKSFSYQALGGHKTSRKPTN
 TSITSGNQELSNNSHNSNSGSVVINVTVNTNGVSQSGKIHTCSICFKSFASGQALGGHKRCH
 YDGGNNNGNGNGSSNSVELVAGSDVDNERWSEESAIGGHRGFDLNLPADQVSVTTS

OTHER GENES IN EVALUATION

SEQ ID NO 36: gi_12698881_ref_AF_332876.1_mRNA, 2xC2H2,
Oryza sativa

AATTCCGGCACGAGGCCACACAGCAACCAGCCAGCTGCCACACTAGCTTGAGGCGAGCGAGCG
 AAGCTTAGCTAGCGGATAGAACACAAGTCGTGATCTGCTGCTTGTGAATTGCGGTGG
 AAGCATGTCAGCGCGTCGTCCATGGAAGCGCTCCACGCCGCGGTGCTCAAGGAGGAGCAGC
 AGCAGCACGAGGTGGAGGAGGCGACGGTCGTGACGAGCAGCAGCGCCACGAGCAGGGAGGAG
 GGCAGCACCTGCCCAAGGGTGGCGAAGCGGAAGCGGTGCGGCCAGCGATCGGAGGA
 GGAGAACCTCGCGCTCTGCTCCTCATGCTCGCCCGCGGCCACCACCGCGTCCAGGCGC
 CGCCTCCGCTCTCGGCTCGGCCCGCCGGCAGGTGCGGAGTTCAAGTGCTCCGTC
 GGCAAGTCCTCAGCTCCTACAGGGCCTCGGCCACAAGACGAGCCACCGGGTCAAGCT
 GCGACTCCGCCCGCAGCTCCGCTTGGCTCCGCCCGTGCCTGCTGCCTCCG
 CCGAGGACCGCGAGCCAGCAGTCATCCACCGCCGCTCTCCGACGGCATGACCAACAGA
 GTCCACAGGTGTTCCATCTGCCAGAAGGAGTTCCCCACCGGGCAGGCCTGGCGGGCACAA
 GAGGAAGACTACGACGGTGGCGTAGGCGCCGGCGCAGTCCACCGCCACCCGGCG
 CCACGGTGGCCGCCAGTCCGAGGTGGGAAGCTCCGCCACGGCCAGTCCGCCACCCGGCG
 TTCGACCTAACCTCCGGCGTGCCTGGAGTTCTGTGTGGCGCCGTGCTCCAAGGGCAAGAA
 GATGTGGGACGAGGAGGAGGAGGTCCAGAGCCCCCTGCCCTCAAGAAGCCCCGGCTCTCA
 CCGCGTAATTCAAGCAGCTGCACGGATCCGATCCGTCAAGAGTTGTCTAGGGAGTGAATT
 CAGTCGAAACACACTATTCTGTTGATTCTGTTGTGCCGCTATTGTTAATTGTTCTGCTT

FIGURE 3 (continued)

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TTGTACAGAGCAAGCGAGTGATACTAGCCATACATACAGTCATAACAGATATAGGTCTAGCT
 CTTCTTGGTTCTTGTAACTGGAACTGTACTGTATCTTACACTTGTCTTGACA
 GTCATATATTGTAGACCAAAAAAAAAAAAAAA

SEQ ID NO 37: gi_12698882_ref_AAK01713.1, 2xC2H2, *Oryza sativa*
 MSSASSMEALHAAVLKEEQQQHEVEEATVVTSSSATSGEEGGHLPGWAKRKRSSRQRSEEE
 NLALCLLMLARGHHHRVQAPPPLSASAPPAGAEFKCSVCGKSFSYQALGGHKTSRVKLP
 TPPAAPVLAAPVAALLPSAEDREPATSSTAASSDGMTNRHRCSCICQKEFPTGQALGGHKR
 KHYDGGVGAGAGASSTELLATVAAESEVGSSNGQSATRAFDLNLPAVPEFVWRPCSKGKMM
 WDEEEEVQSPLAGFKKPRLLTA

SEQ ID NO 38: gi_6434215_ref_AL132966.1_region 116202 ...
 116729, 2xC2H2, *Arabidopsis thaliana*

ATGAAGAGAGACCGGTCCGATTACGAAGAATCCATGAAGCATATAGACATAGTAGAAAGTCT
 AATGATGTTATCTCGAAGTTCGTGGTCAAACAAATCGATGTAAAGCAATCTACCGGAAGCA
 AAACGAACCATAATAACCACCTCGAATGCAAAACGTGTAACCGGAAATTGATTCTTCCAA
 GCTCTGGAGGTCA TAGAGCTAGCCACAAGAAACCTAACGCTGATCGTTGACCAAGAACAGGT
 GAAGCATCGTAACAAAGAGAATGATATGCATAAGTGTACAATTGCGATCAAATGTTGGGA
 CCGGTCAAGCTCTAGGCGGTACATGAGAAAAGCATAGGACGAGCATGATAACCGGAGCAATCG
 ATTGTCCCTCTGTGGTTATTCCAGACCGGTTTAATCGTTGAGTAGCAGCAAGGAGAT
 CTTGGACTAAATCTAACTCCATTGGAAAATGATCTGTGTTAATCTTGGGAAGAATTGG
 TTCCACAAATTGATTTGAAGTTGTGAATTAG

SEQ ID NO 39: gi_6729511_ref_CAB67667.1, 2xC2H2, *Arabidopsis thaliana*
 MKRDRSDYEESMKHIDIVESLMMILSRSFVVKQIDVKQSTGSKTNHNNHFECKTCNRKFDSFQ
 ALGGHRASHKKPKLIVDQEVKHRNKENDMHKCTICDQMFGTGQALGGHMRKHTSMITEQS
 IVPSVYYSRPVFNRCSSSKEILDNLPLENDLVLIFGKNLVPQIDLKFVN

SEQ ID NO 40: ref_CA279020, 2xC2H2, sugar cane
 CCTAACCAAGCATTAGCTTTCAAATCAACAAGCCTGCCGTGACCGATCGATGGCCATCACC
 CACGACGACTACGTCTCCCTCTGCCATGGCGCTCGCAGCCGCGGGAGGCAGGCGCAAGAGTCCG
 TGGTTAACAAACCGAGTACGCTCTGAACACGGCTGCCGGACAGCGACGGCGCAAGAGTCCG
 AGCTCCGCTTCCGGTCTCGTCTGGCAAGGGCTTCGGTCCGCTCGCACCGGACTGGCGGG
 CACAAGGCCAGCCACCGCAAGCCGACGCTCGTACAGGCACATGCGTCGCTCAGCCGGAGG
 CGCGGCGTCGTCGGTAACAATGACCTCGGCCGTAGGCAGTGGCAGGGAGGCACA
 GGTGCACGGTGTGCCATGGAGCTCGCAGGGNGCAAGCGCTGGCGGGACAAGAGGTGC
 CATTACTGGGACGGGCTCTGGTCTCGCTACCGCGTCGCTCGGCCATGGGGTCCGGT
 GACCGTCAAGGGCTTGATCTGAATTGGTGCCGTGCCGCCGCGATGGCGCCAACGCTG
 CGACAAGGTGGGGAGAGGAGAANNAAGTCANAAACCTGGCGGTCAAGAGAAGGCAGGCTTG
 CCGGTCCGCTTGGACCCTAATTAAACGATTAGAAGTCCTTTTTAATAATTAAAGAGTTC
 TTTGAAGAAGGTTGTAAGTTTCAACCTGTTCTTAATGGATTGGGTGCTGGCGAA
 ATTTAAAATGGATTAAATTGCGCTCACTCTTTTTTATTACACCCTTTTT
 TTTTAGAAGAAGAAGA

FIGURE 3 (continued)

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SEQ ID NO 41: gi_18027011_ref_AF254447.1, 2xC2H2, *Arabidopsis thaliana*

TTCCCTTCTCTCCTCTCTCTCTCACCATGACTGATCCTTATTCCAATTCTTCACA
 GACTGGTCAAGTCTAATCCTTTCACCATCACCTAATTCCCTCCACTAACCCCTCTCCTCA
 TCCTCTCCTCCTGTTACTCCTCCCTCTCCTCTTCTCCCTCAATCCGGAGACCTCC
 GCCGTCCACCGGCCACCAACTCCTCCTCCTCTCCCTCCGAGAACGCCCTCCCTCTC
 CTCAGCCTCAGCCCCGCCAACAAACAAGACCCATCACACCACATGACCACCTTATTCA
 AGAACACCTTCACACCTCCATGGATGTCGACTACGATCATCACCATCAAGATGATCATCATA
 ACCTCGATGACGATGACCATGACGTACCGTTGCTCTCACATAGGCCTCCAAGCCCTAGT
 GCTCAAGAGATGGCCTTTGCTCATGATGTCCTCTCCTCTCGAGGGACCACTCA
 TCATCAGGAGACATGAATACAAGAAAGACCTCGACCATGAGTACAGCCACGGAGCTGTCG
 GAGGAGGAGAAGATGACGATGAAGATCAGTCGGCGGAGACGGCGCTGAGAATCAGCAGA
 CTCACAAGGGTCAATATTGGATCCCTACACCTCTCAGATTCTCATTGGCCTACTCAGTT
 CTCATGTCCTGTTGCTTCAAAACCTCAACAGATAACAATAACATGCAGATGCATATGTGG
 GACATGGATCACAATACAGAAAAGGACCTGAATCTCTAAGGGAAACACAACAGGAATG
 CTAAGGCTTCCGTGCTATTGCTGCGCCCCAGGCTGTCGAACAAACATTGACCATCCAAGGGC
 AAAGCCTCTCAAAGACTTCAGAACCCCTCAAACACATTACAAGAGAAAACATGGATCAAAC
 CTTTCATGTTGAGGAAATGTGAAAGGCTTCGCAGTCGGAGGGACTGGAGAACATGAG
 AAGAATTGTGGAAACTTGGTATTGCAATATGTGGATCTGATTCAAGCACAAGAGATCTCT
 CAAAGATCACATCAAGGCTTTGGGAATGGTCAATGGAGCCTACGGAATTGATGGGTTGATG
 AAGAAGATGAGCCTGCCTCTGAGGTAGAACATTAGACAATGATCATGAGTCATGCAGTCT
 AAATAGCTTATATATATTACTATAAGTACTAAGTAATTGGTATATATATTAAATTATAAGAA
 ACCTAAATCTATGGACCAAGTTGATGGAGGTAGGGCTTTCAAACCTAAAGCTATATCAT
 CTAATTGATCATAGGAAAAAAATGAATCAAGAGCACTTGGAAAATTAAATTGTATCTTA
 GCTTCCTAGTTAAATTATTGCAAGACAATGTAGCAGTCAACCAATGAGGTTCCAACGGT
 TTATTCTATTGTATATTATTGTCATTAGCTCACCTTCGTTAATTGCAAGGACATAA
 CTTATAATGTTAAATTATG

SEQ ID NO 42: At3g57670, 2xC2H2, *Arabidopsis thaliana*

MTDPYSNFFTDFWKSNPFHYPNSSTNPSPHPLPPVTPPSSFFFFPQSGDLRRPPPPPTPPP
 SPPLREALPLLSLSPANKQDHHHNHDHLIQEPPSTSMDVDYDHHDHQDDHHNLDDDDHDVTV
 ALHIGLPSPSAQEMASLLMMSSSSSSRTTHHEDMNHKKLDHEYSHGAVGGGEDDDDEDSV
 GGDGGCRISRLNKQYWIPTPSQILIGPTFSCPVCFKTFNRYNNQMHWGHGSQYRKGPE
 SLRGQTGMLRLPCYCCAPGCRNNIDHPRAKPLKFRTLQTHYKRKHGIKPFMCRKCGKAF
 AVRGDWRTHEKNCGLWYCICGSDFKHKRSLKDHIAFGNGHGAYGIDGFDEEDEPASEVEQ
 LDNDHESMQSK

SEQ ID NO 43: gi_18676370_ref_AJ311810.2, 2xC2H2, *Arabidopsis thaliana*

ATCTACACACTACTCACATCTCATCTCTAGCACATACCCATCAAACCATATAGAT
 ACGGTGCTTTATTCTTGATCTCTCTTCTTGTCTCTCAGAGTCATGCTAAT
 CCAGCTGTTCGAATCTCAACAAATGGATGTGACCATATAAGCTCAACTATTCCACTTC
 TCTCTTACATTACAACACTCAGGTAGCTACTATTACTCTAATACCAAAACCCCTAATT
 ACATTAATCATACTCATACCACTTCCACTTCCCTAACTCACCCCCACTAAAGAGAACGCTTT
 CCTCTCTTAGCTTAAGCCCCATAAGGCACCAAGAACACAAGACCAACACTATTGATGGA
 CACCCATCAAATTAGCTCTCAAACCTTGATGATCCTCTTGTGACTGTGGATCTCATH
 TAGGGTTACCAAACACTACGGTGTGGTGAGAGCATTAGGAGCAATATTGCTCCTGATGCAACC

FIGURE 3 (continued)

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ACGGACGAGCAAGATCAAGATCATGACCGAGGAGTAGAAGTCACAGTTGAGTCCCACCTTGA
 TGATGATGATGATCATCATGGAGATCTACACAGAGGTACACTATTGGATTCCCTACTCCTT
 CTCAGATTGATTGGTCTACACAGTCACTTGTCCTTTGCTTCAAGACATTCAACAGA
 TACAACAACATGCAGATGCACATGTGGGACACGGCTACAATACAGAAAGGGACCAGAAC
 CTTAAGAGGAACCCAACCAACAGGAATGCTAAGACTACCATGTTCTGCTGTGCACCCGGTT
 GCAAGAACAAACATTGACCACCCACGAGCCAAGCCTCTAAGGACTTTCGAACCCCTCAAACA
 CATTACAAACGTAACATGGGTCTAAACCATTGCTGCTATGTGTGGTAAGGCCTTTGC
 AGTGAAGGAGATTGGAGAACGCACTGAGAAGAATTGTGAAAGCTTGGTATGCTCTGTG
 GCTCGGATTAAAGCACAAGAGGTGCTTAAGGACCATGTCAAGGCCTTGGAAATGGTCAT
 GTTCCTGTTGGATTGATAGTTGGAGGAGATCATGAGGACTACTATGATGCTGCTCTGA
 TATCGAGCAATAAGATGATAGCAACAACAATGAGTGTAAATTAGGGTTTGTATTTC
 CTCTCATGCATTAGTTGATTGATGACGTGTTAGTTGGTCTTCGGATCTTGT
 TATTTGTTTGAGCTGTTTTTTAATTACTAAGAAGTTAATTATCATCTAAAGATT
 C

SEQ ID NO 44: gi_18376498_ref_CAC86167.1, 2xC2H2, *Arabidopsis thaliana*

MSNPACSNLFNNNGCDHNSFNYSTSLSYIYNHSYYSNNTNPYINHHTTSTSPNSPPLR
 EALPLLSLSPIRHQEQQDQHYFMDTHQISSLNFDDPLVTVLHLGLPNYGVGESIRSNIAP
 DATTDEQDQDHDRGVEVTVESLHDDDDDHGDLHGRHHYWIPTPSQILIGPTQFTCPLCFKT
 FNRYNNNMQMHWGHSQYRKGPESLRGTQPTGMLRLPCFCCAPGCKNNIDHPRAKPLKDFRT
 LQTHYKRKHGSKPFACRMCGKAFAVKGDWRTHEKNCGLWYCSCGSDFKHKRSLKDHVKAFG
 NGHVPCGIDSFGGDHEDYYDAASDIEQ

SEQ ID NO 45: gi_7798991_ref_AL355775.1_region 7957 ... 8451,
 2xC2H2, *Arabidopsis thaliana*

ATGGTTGCGAGAAGTGAGATAGTGGAGATAACGGCGCGAAATGTTGATGTT
 GTTATCAAGAGTTGGAGAATGCGCGGAGGAGAGAACGAGTTTCCGATGCAAGACTT
 GTCTTAAAGAGTTTCGTCGTTCAAGCTTGGAGGTACATCGTCAAGCCACAAGAAACTC
 ATTAACAGTAGCGATCCATCACTTCTGGATCCTGTCTAACAAAGAAAACGAC
 GTCTCATCCTGTCGATATGTGGCGTGGAGTTCCGATGGGCAAGCTCTGGTGGTCACA
 TGAGGAGACATAGGAGTGAGAAAGCCTCACCAAGGCACGTTGGTACACGTTTTACCG
 GAGACGACGACGGTGACGACTTGAAAAATCGAGTAGTGGGAAGAGAGTGGCTTGG
 CTTAGATTGAGAGTTAGTCAATTGGAAGTTGGAGTTGGGAAGAACGATTCTGA

SEQ ID NO 46: gi_7798996_ref_CAB90935.1, 2xC2H2, *Arabidopsis thaliana*

MVARSEEVEIVEDTAAKCLMLLSRVGECGGGGKRVFRCKTCLKEFSSFQALGGHRASHKKL
 INSSDPSLLGSLSNKKTATSHPCPICGVEMQALGGHMRHRSEKASPGTLVTRSFLP
 ETTTVTTLKSSSGKRVACLDLDSMESLVNWKLELGRITIS

SEQ ID NO 47: gi_9755794_ref_AL391143.1_region 31730 ...
 32938, 2xC2H2, *Arabidopsis thaliana*

ATGGAAGACGAACATCAAGATCTCCATAAACCCATTAAATGGAGCTTGCAGACCTCAAGAT
 TACTCGGTACAGAAAGAACAGAAAAGTCTACGAACCAACAGCAAGATGTACTTGTACT
 ATGGTCTAAGGGAAAACTCGAAGAAGAAAACCCAGGAATCTCCGGAACCAATGAAGAAGATT
 TTGTTTCGATGCCAAGAATGTGGAAAAGGGTTCCGGTACGAGAAATTTAAGAATCATCG

FIGURE 3 (continued)

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CTCGATGATGCATTATGCCGAACGAGAAGGTTGTGAAGAATCCTGATGACTCTGCTCT
 GTAGCCTGGGTTGTGAAGAAGAAAAGATCAAGACTTGGTAGGTCTGGGAAGACTTTA
 TTTACTACGTTCTGAACCGAGTTCTATTGATGCGACTGATGAAGAATTAGAAGTGGC
 GGATTGTTGATTCTATTGCTAAGAGTGTCTCCAAGGTTGTAGACGAATTGAAAAGTCTT
 CTGAGGCAGTACGTACTCCTGAAACACCTGAAAGTAGCTATGATTTGGGTGTTGCTC
 AACAAAGAAACCGAGAAAAGGTTGAATTGGAATCTGGGTTTAAGTAATGAGCAAAGACT
 TATGGAAGAAGGGTTAGTAGTTATGGAACATCGAAAGAACAGCTAGCTTCTGAGAGACG
 AAAACAGATTGGATCAGCAGAAACGGAGAAAAGATGGTGAATTGAAATCCGGACTTTGAGT
 AATGAGCAAAGACTGCTAGAAGAAGAGATTACTACTCCTGACATCAAAGGTCCAGCGAG
 TTCCTTGAGACACAAGTGTGCTTGGATCGAAATGGAGGTGAATTGGTCTGAGTTTGA
 GTAATGAGCAAACACTGATGGAAGAAACATGGAAGAACAGCTGGCAGTTGAATCTAGGTTTACAGAAT
 CATGAATTGATCAGCGAAAATGCGAGAAAGCTGGCAGTTGAATCTAGGTTTACAGAAT
 TGAGCTGGAGTAGGAGCTATGGAGTACTTCTCAGATACTGATATGCTACGCAATCTG
 ATAAGAAGAACGTTGAGCATCGATGCGAGTTGTGCAACAAGATATTCTGCTTATCAAGCT
 CTAGGGGGTCACTCAGACGTTCATCGGATGAGCAAATGTAAGAACAGAACATGGCATAGA
 GGAATCAGTTGAACCCAGGATGACTCTGTGA

SEQ ID NO 48: gi_9755803_ref_CAC01747.1, 2xC2H2, Arabidopsis thaliana

MEDEHQDLHKPINGALRDLKIRSQKETEKSTNQQODVTCYYGLRENSKKKTQESPEPMKKI
 LFRCEECGKFRYEKYFKNHRSMHLSPNEKVCEESLMTSLRGVKKKRSRLGRSGKTL
 FTTFLEPSSI FDATDEELEVADCLILSKSAPKVVDELKSLSEAVRVTPETPESSYDLGCLL
 NKKPRKGGELESVLSNEQRIMEEGFSYGTSKEPASFLRDENRLDQQKRRKDGEFESGLLS
 NEQRILEEEITTPVTFKGPASSLRHKCALDRNGGEFGPEFLSNEQTLMEETWKEPVSFLEDK
 HEFDQRKMREAGDFESRFYRIELGVGAMECTSSDTDMLTQSDKKNVEHRCRLCNKIFSSYQA
 LGGHQTFHRMSKCKNKKNGIEEVEPRMTL

SEQ ID NO 49: gi_1418338_ref_X98678.1, 2xC2H2, Arabidopsis thaliana

CTTGTAGTTCACTCCACATAATAACACAAAGATTCATTCTCTTCTCCATAATTCGAA
 GTTCTTGAAATTGGGTTTGTCTTGATTGTTCTGATTGGTTGGTCTTCTTTCT
 TACTATATTGGATATGATGGTCAAGATGAGGTTGGGAGTGTACAGACGCAAATCATA
 AAAGGGAAACGTACGAAGCGACAAAGATCGTCTCGACGTTGTGGACGGCGGCGACAAC
 AGTGAATTCAACAAGTTCATCGGCCGGTGGAAAGTGGAGGAGAAAGAGACTGTTAGATGAAT
 ACAACTCGCCGGTTCTGTCCTCGTACTGATTGTACGCAAGAAGAACATGGCG
 ATTTGTCATCATGTTAGCTCGTGGGACAGTCTTCCATGCCGGATCTCAAGAACTCGAG
 AAAAATCATCAGAAGATTCTGTCGGAGAATTCTAGTTCTATGTGTACGAGTGTAAACGT
 GTAACCGGACGTTCTGTCGGTCAAGCACTGGTGGACACAGAGCGAGCCACAAGAACCG
 AGGACGTCGACTGAGGAAAGACTAGACTACCCCTGACGCAACCCAAAGTCTAGTCAGTGCATCAGA
 AGAAGGGAAAACAGTCATTCAAAGTTCCGGCTCAGCCCTAGCTTACAGGGCAAGTAACA
 TCATCAACAAGGAAACAAAGTACACGAGTGTCCATCTGCGGTTCTGAGTTCACTCCGGG
 CAAGCTCTCGGTGGTCACATGAGGCCGACAGGACAGCCGTAACACGATTAGCCCCGTTGC
 AGCCACCGCAGAGTAAGCAGAACAGTACAGAGGAAGAGATTGAGATCAATATAGGCCGTT
 CGATGGAACAGCAGAGGAATATCTACCGTTGGATCTTAATCTACAGCACCCAGGAGATGAT
 CTAAGAGAGTCCAAGTTCAAGGGATAGTATTCTCAGCAACACCAGCGTTAATAGATTGTCA
 TTACTAGTTGTTTTTACTACATAATGATGAAATATTGTGAATTCTCTTACTTACT
 ACTATATTGTTGATCAAAAAAAAAAAAAAA

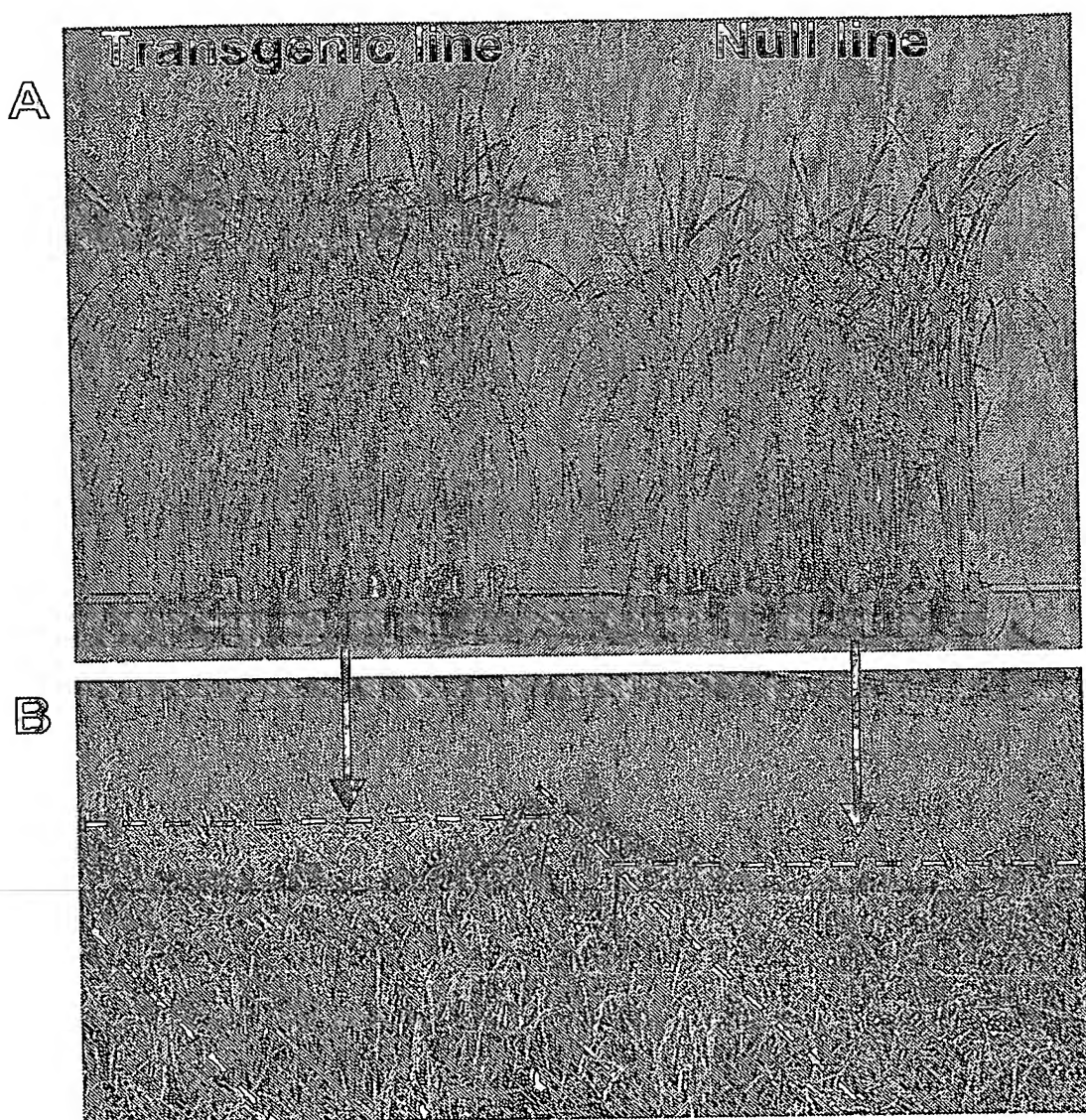
FIGURE 3 (continued)

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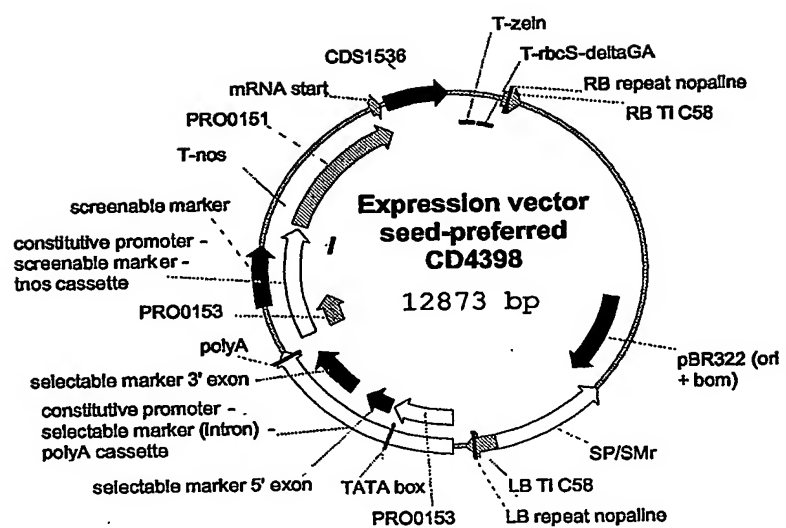
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FIGURE 3 (continued)

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**FIGURE 4**

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**FIGURE 5**

SEQUENCE LISTING

<110> CropDesign N.V.

<120> Plants having modified growth characteristics and a method for
making the same

<130> CD-070-PCT

<160> 50

<170> PatentIn version 3.1

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<211> 692

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															25
															30

Lys	Gly	Lys	Arg	Ser	Lys	Arg	Ser	Arg	Ser	Asp	Phe	His	His	Gln	Asn
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															40
															45

Leu	Thr	Glu	Glu	Tyr	Leu	Ala	Phe	Cys	Leu	Met	Leu	Leu	Ala	Arg	
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															55
															60

Asp	Asn	Arg	Gln	Pro	Pro	Pro	Pro	Pro	Ala	Val	Glu	Lys	Leu	Ser	Tyr
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															70
															75
															80

Lys	Cys	Ser	Val	Cys	Asp	Lys	Thr	Phe	Ser	Ser	Tyr	Gln	Ala	Leu	Gly
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															90
															95

Gly	His	Lys	Ala	Ser	His	Arg	Lys	Asn	Leu	Ser	Gln	Thr	Leu	Ser	Gly
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Gly Asn Asn Asn Ile Asn Thr Ser Ser Val Ser Asn Ser Glu Gly Ala		
165	170	175
Gly Ser Thr Ser His Val Ser Ser His Arg Gly Phe Asp Leu Asn		
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Ile Pro Pro Ile Pro Glu Phe Ser Met Val Asn Gly Asp Asp Glu Val		
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<213> Artificial sequence

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<220>
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<210> 8
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<223> Ser can be serine or no amino acid

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<223> Xaa can be any amino acid

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<212> DNA
<213> Datisca glomerata

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gacgacccca gcttgaatta ccttgagccca tggaccaagc gtaagcgttc caagcgtacg 180
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cgtcttatca ggcttgggt gggcacaagg ccagccacag aaagctcgat ggcggcgaag 420
atcaatcgac ttcccttgcc accacgaatt cagccaccgt cactaccacc acagcctccg 480

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accacaacaa taccaccaac acggaaagca acgggtggcat gagcatgacc tccgaagtag	660
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attaatatat ttctgttaca taaatttgcgatccggatccggatccggatccggatccggat	960
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<211> 247
<212> PRT
<213> *Datisca glomerata*

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Lys Arg Ser Lys Arg Thr Arg Leu Asp Ser Pro His Thr Glu Glu Glu	
35 40 45	
Tyr Leu Ala Phe Cys Leu Ile Met Leu Ala Arg Gly Arg Val Ala Ser	
50 55 60	
Ala Asn Arg Arg Asp Ser Gln Ser Ser Ile Gln Ile Gln Pro Glu Ala	
65 70 75 80	
Thr Thr Ser Ala Thr Lys Val Ser Tyr Lys Cys Ser Val Cys Asp Lys	
85 90 95	
Ala Phe Ser Ser Tyr Gln Ala Leu Gly Gly His Lys Ala Ser His Arg	
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Lys Leu Ala Gly Gly Glu Asp Gln Ser Thr Ser Phe Ala Thr Thr Asn	
115 120 125	
Ser Ala Thr Val Thr Thr Ala Ser Gly Gly Gly Arg Ser	
130 135 140	
His Glu Cys Ser Ile Cys His Lys Ser Phe Pro Thr Gly Gln Ala Leu	
145 150 155 160	
Gly Gly His Lys Arg Cys His Tyr Glu Gly Ser Ile Gly Gly Asn Ser	
165 170 175	
Ile His His His Asn Asn Thr Thr Asn Ser Gly Ser Asn Gly Gly Met	
180 185 190	
Ser Met Thr Ser Glu Val Gly Ser Thr His Thr Val Ser His Ser His	
195 200 205	
Arg Asp Phe Asp Leu Asn Ile Pro Ala Leu Pro Glu Phe Arg Ser Asn	
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Lys Pro Arg Ile Leu Met Lys
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 aatttcaatg aactcgttga attttttagt tattttcga ctatataattt tggagaattt 840
 tgagagttaac tataatttga ttttgtacat agtacttgg 900
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 <211> 240
 <212> PRT
 <213> Glycine max

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Arg Ser Arg Asp His Pro Ser Glu Glu Glu Tyr Leu Ala Leu Cys Leu
 35 40 45

Ile Met Leu Ala Arg Gly Gly Thr Thr Val Asn Asn Arg His Val
 50 55 60

Ser Pro Pro Pro Leu Gln Pro Gln Pro Thr Pro Asp Pro Ser
 65 70 75 80

Thr Lys Leu Ser Tyr Lys Cys Ser Val Cys Asp Lys Ser Phe Pro Ser
 85 90 95

Tyr Gln Ala Leu Gly Gly His Lys Ala Ser His Arg Lys Leu Ala Gly
 100 105 110

Ala Ala Glu Asp Gln Pro Pro Ser Thr Thr Ser Ser Ala Ala Ala

115

120

125

Thr Ser Ser Ala Ser Gly Gly Lys Ala His Glu Cys Ser Ile Cys His
 130 135 140

Lys Ser Phe Pro Thr Gly Gln Ala Leu Gly Gly His Lys Arg Cys His
 145 150 155 160

Tyr Glu Gly Asn Gly Asn Asn Asn Asn Ser Asn Ser Val Val
 165 170 175

Thr Val Ala Ser Glu Gly Val Gly Ser Thr His Thr Val Ser His Gly
 180 185 190

His His Arg Asp Phe Asp Leu Asn Ile Pro Ala Phe Pro Asp Phe Ser
 195 200 205

Thr Lys Val Gly Glu Asp Glu Val Glu Ser Pro His Pro Val Met Lys
 210 215 220

Lys Pro Arg Leu Phe Val Ile Pro Lys Ile Glu Ile Pro Gln Phe Gln
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<210> 14

<211> 1006

<212> DNA

<213> *Medicago sativa*

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<212> PRT

<213> *Medicago sativa*

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 35 40 45

Glu Glu Tyr Leu Ala Leu Cys Leu Ile Met Leu Ala Arg Ser Gly Asn
 50 55 60

Asn Asn Asp Lys Lys Ser Asp Ser Val Ala Thr Pro Leu Thr Thr Val
 65 70 75 80

Lys Leu Ser His Lys Cys Ser Val Cys Asn Lys Ala Phe Ser Ser Tyr
 85 90 95

Gln Ala Leu Gly Gly His Lys Ala Ser His Arg Lys Ala Val Met Ser
 100 105 110

Ala Thr Thr Ala Glu Asp Gln Ile Thr Thr Thr Ser Ser Ala Val Thr
 115 120 125

Thr Ser Ser Ala Ser Asn Gly Lys Asn Lys Thr His Glu Cys Ser Ile
 130 135 140

Cys His Lys Ser Phe Pro Thr Gly Gln Ala Leu Gly Gly His Lys Arg
 145 150 155 160

Cys His Tyr Glu Gly Ser Val Gly Ala Gly Ala Gly Ser Asn
 165 170 175

Ala Val Thr Ala Ser Glu Gly Val Gly Leu Ser His Ser His His Arg
 180 185 190

Asp Phe Asp Leu Asn Leu Pro Ala Phe Pro Asp Phe Ser Lys Lys Phe
 195 200 205

Phe Val Asp Asp Glu Val Phe Ser Pro Leu Pro Ala Ala Lys Lys Pro
 210 215 220

Cys Leu Phe Lys Leu Glu Ile Pro Ser His Tyr
 225 230 235

<210> 16
 <211> 1061
 <212> DNA
 <213> Nicotiana tabacum

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gtaaaatttgg	tcatgtgatt	ttatTTTtag	gaaaaggaat	tattgtattgt	tttaccggtt	960
tattctttagg	gtggatttat	gtacaggggag	tgaatcatttc	attgggtttta	cactttctta	1020
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<212> PRT
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 20 25 30

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35 40 45

Glu Glu Glu Tyr Leu Ala Leu Cys Leu Ile Met Leu Ala Arg Ser Gly
50 55 60

Thr Gly Thr Arg Thr Gly Leu Thr Asp Ala Thr Thr Ser Gln Gln Pro
65 70 75 80

Ala Asp Lys Lys Thr Ala Glu Leu Pro Pro Val His Lys Lys Glu Val
85 90 95

Ala Thr Glu Gln Ala Glu Gln Ser Tyr Lys Cys Ser Val Cys Asp Lys
 100 105 110

Ala Phe Ser Ser Tyr Gln Ala Leu Gly Gly His Lys Ala Ser His Arg
 115 120 125

Lys Thr Thr Thr Ala Thr Ala Ala Ser Asp Asp Asn Asn Pro Ser
 130 135 140

Thr	Ser	Thr	Ser	Thr	Gly	Ala	Val	Asn	Ile	Ser	Ala	Leu	Asn	Pro	Thr
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Gly Arg Ser His Val Cys Ser Ile Cys His Lys Ala Phe Pro Thr Gly
 165 170 175

Gln Ala Leu Gly Gly His Lys Arg Arg His Tyr Glu Gly Lys Leu Gly
180 185 190

Gly Asn Ser Arg Asp Leu Gly Gly Gly Gly Gly Gly Gly His Ser Gly
195 200 205

Ser Val Leu Thr Thr Ser Asp Gly Gly Ala Ser Thr His Thr Leu Arg
210 215 220

Asp Phe Asp Leu Asn Met Pro Ala Ser Pro Glu Leu Gln Leu Gly Leu
225 230 235 240

Ser Ile Asp Cys Gly Arg Lys Ser Gln Leu Leu Pro Met Val Gln Glu
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Val Glu Ser Pro Met Pro Ala Lys Lys Pro Arg Leu Leu Phe Ser Leu
 260 265 270

Gly

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<211> 1213

<212> DNA

<213> Oryza sativa

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 aaaaaaaaaaaa aaa 1213

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<212> PRT

<213> Oryza sativa

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Ser Ser Ala Thr Ser Gly Glu Glu Gly Gly His Leu Pro Gln Gly Trp
 35 40 45

Ala Lys Arg Lys Arg Ser Arg Arg Gln Arg Ser Glu Glu Glu Asn Leu
 50 55 60

Ala Leu Cys Leu Leu Met Leu Ala Arg Gly Gly His His Arg Val Gln
 65 70 75 80

Ala Pro Pro Pro Leu Ser Ala Ser Ala Pro Pro Pro Ala Gly Ala Glu
 85 90 95

Phe Lys Cys Ser Val Cys Gly Lys Ser Phe Ser Ser Tyr Gln Ala Leu
 100 105 110

Gly Gly His Lys Thr Ser His Arg Val Lys Leu Pro Thr Pro Pro Ala
 115 120 125

Ala Pro Val Leu Ala Pro Ala Pro Val Ala Ala Leu Leu Pro Ser Ala
 130 135 140

Glu Asp Arg Glu Pro Ala Thr Ser Ser Thr Ala Ala Ser Ser Asp Gly
 145 150 155 160

Met Thr Asn Arg Val His Arg Cys Ser Ile Cys Gln Lys Glu Phe Pro
 165 170 175

Thr Gly Gln Ala Leu Gly Gly His Lys Arg Lys His Tyr Asp Gly Gly
 180 185 190

Val Gly Ala Gly Ala Gly Ala Ser Ser Thr Glu Leu Leu Ala Thr Val
 195 200 205

Ala Ala Glu Ser Glu Val Gly Ser Ser Gly Asn Gly Gln Ser Ala Thr
 210 215 220

Arg Ala Phe Asp Leu Asn Leu Pro Ala Val Pro Glu Phe Val Trp Arg
 225 230 235 240

Pro Cys Ser Lys Gly Lys Lys Met Trp Asp Glu Glu Glu Val Gln
 245 250 255

Ser Pro Leu Ala Phe Lys Lys Pro Arg Leu Leu Thr Ala
 260 265

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 <211> 1020
 <212> DNA
 <213> Petunia x hybrida

<400> 20
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 aatacaaaca agaaaatttt ctctctatac ttgattgagt tttagtaaggc aaacaagaaa 180
 actatcatgg cacttgaagc attgaattct ccaactacaa caacaccacc atcattccaa 240
 tttgagaaca acgggcttaa gtaccttgag agttggacaa aaggtaaaag atcaaaaagg 300
 caacgcagca tggAACgaca gtgtactgaa gaagagtatt tagcactttg tcttatcatg 360
 ctagcactgtc gcgatggttc tgtaataac tcacgtctc taccaccacc accactacca 420
 ccatcagttc cagtaacgtc gcaataaaac gcgacgttat tggAACagaa gaatttgcac 480
 aagtgttccg tttgtggtaa agggtttggc tcttatacaag ctttaggtgg acataaagca 540
 agtcacccga aacttgtcag catgggagga gatgaacaaat ctactacttc cactact 600
 aacgtaaacgg gaaactgttcc cgctaacgtt aacggtaacgg gaaagaactca cgaatgtca 660
 atttgtcaca agtgctttcc tactggacaa gcttttaggtg gtcataaaaag gtgccactat 720
 gacgggtggta acggtaacgg taacggaaat gtaagtgtt gggtgacgtc atctgaaggt 780
 gtgggggtcca ctattagtcg tcaccgtgac tttgacttga atattcccgc gttgcccggag 840
 ttttggccgg gatttggttc cggcgaggat gaggtggaga gtcctcatcc agcaaagaag 900

tcaaggctat ctcttccacc taaacttcaa ttattcaaag gattatagag ggaatattga 960
tttgcacag gaagatttat taggattcac gaatttttg ttgactagtt tatgtaatat 1020

<210> 21
<211> 253
<212> PRT
<213> Petunia x hybrida

<400> 21
Met Ala Leu Glu Ala Leu Asn Ser Pro Thr Thr Thr Thr Pro Pro Ser
1 5 10 15

Phe Gln Phe Glu Asn Asn Gly Leu Lys Tyr Leu Glu Ser Trp Thr Lys
20 25 30

Gly Lys Arg Ser Lys Arg Gln Arg Ser Met Glu Arg Gln Cys Thr Glu
35 40 45

Glu Glu Tyr Leu Ala Leu Cys Leu Ile Met Leu Ala Arg Ser Asp Gly
50 55 60

Ser Val Asn Asn Ser Arg Ser Leu Pro Pro Pro Pro Leu Pro Pro Ser
65 70 75 80

Val Pro Val Thr Ser Gln Ile Asn Ala Thr Leu Leu Glu Gln Lys Asn
85 90 95

Leu Tyr Lys Cys Ser Val Cys Gly Lys Gly Phe Gly Ser Tyr Gln Ala
100 105 110

Leu Gly Gly His Lys Ala Ser His Arg Lys Leu Val Ser Met Gly Gly
115 120 125

Asp Glu Gln Ser Thr Thr Ser Thr Thr Thr Asn Val Thr Gly Thr Ser
130 135 140

Ser Ala Asn Val Asn Gly Asn Gly Arg Thr His Glu Cys Ser Ile Cys
145 150 155 160

His Lys Cys Phe Pro Thr Gly Gln Ala Leu Gly Gly His Lys Arg Cys
165 170 175

His Tyr Asp Gly Gly Asn Gly Asn Gly Ser Val Ser Val Gly
180 185 190

Val Thr Ser Ser Glu Gly Val Gly Ser Thr Ile Ser His His Arg Asp
195 200 205

Phe Asp Leu Asn Ile Pro Ala Leu Pro Glu Phe Trp Pro Gly Phe Gly
210 215 220

Ser Gly Glu Asp Glu Val Glu Ser Pro His Pro Ala Lys Lys Ser Arg
225 230 235 240

Leu Ser Leu Pro Pro Lys Leu Glu Leu Phe Lys Gly Leu
245 250

<210> 22
 <211> 786
 <212> DNA
 <213> *Triticum aestivum*

<400> 22
 atgtcgtcgatggccatggaa agcgctccac gcccgtatcc cggagcagca ccagctggac 60
 gttgaggcgg ctgcggctgt cagcagcgcc accagccgcg aggagagcgg ccacgtgctg 120
 caggggtggg ccaagaggaa gcgatcgcc cgccagcgct cegaggagga gaacctcgcg 180
 ctctgcctcc tcatgtctc gcgcggcgcc aagcagcgta ttcaggcgcc gcagccggag 240
 tcgttcgtcgcc tcccgaggatc aagtgtcccg tctgcggcaa gtcccttcagc 300
 tcctaccagg cgctcggagg ccacaagacg agccacccggg tgaagcagcc gtctccccc 360
 tctgtatggccg ctgtcgcccc actcggtggcc ctcccgcccg tcggccat cctggcgcc 420
 gccgagccgg ccacgtcgatcc caccggcgcc tcctccgacg gcgcgacaa cagagtccac 480
 aggtgttcca tctgcggaaaaa ggatgtcccg actggggcagg cgctcgccgg gcacaagagg 540
 aagcaactacg acggaggcgt gggcgccgcg gcctcgatcg ccgagcttct ggccggcccg 600
 gccgcccggat ctgagggtggg gaggcaccggc aacggggatc cccggcccg ggcccttcgac 660
 ctgaacattc cggccgtgccc ggagttcgatc tggaggccgt gcgcacaaggg caagatgtatc 720
 tggaggacg atgaggaggt gcagagcccc ctcgccttca agaagcctcg gcttctcacc 780
 gcttga 786

<210> 23
 <211> 261
 <212> PRT
 <213> *Triticum aestivum*

<400> 23
 Met Ser Ser Ser Ala Met Glu Ala Leu His Ala Leu Ile Pro Glu Gln
 1 5 10 15
 His Gln Leu Asp Val Glu Ala Ala Ala Val Ser Ser Ala Thr Ser
 20 25 30
 Gly Glu Glu Ser Gly His Val Leu Gln Gly Trp Ala Lys Arg Lys Arg
 35 40 45
 Ser Arg Arg Gln Arg Ser Glu Glu Glu Asn Leu Ala Leu Cys Leu Leu
 50 55 60
 Met Leu Ser Arg Gly Gly Lys Gln Arg Val Gln Ala Pro Gln Pro Glu
 65 70 75 80
 Ser Phe Ala Ala Pro Val Pro Ala Glu Phe Lys Cys Ser Val Cys Gly
 85 90 95
 Lys Ser Phe Ser Ser Tyr Gln Ala Leu Gly Gly His Lys Thr Ser His
 100 105 110
 Arg Val Lys Gln Pro Ser Pro Ser Asp Ala Ala Ala Pro Leu
 115 120 125
 Val Ala Leu Pro Ala Val Ala Ala Ile Leu Pro Ser Ala Glu Pro Ala
 130 135 140
 Thr Ser Ser Thr Ala Ala Ser Ser Asp Gly Ala Thr Asn Arg Val His
 145 150 155 160
 Arg Cys Ser Ile Cys Gln Lys Glu Phe Pro Thr Gly Gln Ala Leu Gly

165

170

175

Gly His Lys Arg Lys His Tyr Asp Gly Gly Val Gly Ala Ala Ala Ser
 180 185 190

Ser Thr Glu Leu Leu Ala Ala Ala Ala Glu Ser Glu Val Gly Ser
 195 200 205

Thr Gly Asn Gly Ser Ser Ala Ala Arg Ala Phe Asp Leu Asn Ile Pro
 210 215 220

Ala Val Pro Glu Phe Val Trp Arg Pro Cys Ala Lys Gly Lys Met Met
 225 230 235 240

Trp Glu Asp Asp Glu Glu Val Gln Ser Pro Leu Ala Phe Lys Lys Pro
 245 250 255

Arg Leu Leu Thr Ala
 260

<210> 24

<211> 1026

<212> DNA

<213> Capsicum annum

<400> 24

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 aacggatatac gatatggcac ttgaagcttt gaattctcca actggcacac caactccgccc
 accgtttcaa tttgagagcg acggccaaaca gcttcgatata atcgaaaact ggaggaaggg
 aaagagatct aaaaggtcac gcagcatgga gcaccagcct actgaggaag aataacttagc
 gctttgttg atcatgcttg cacgtacggg tggctccgtt aatcatcaac gatctctacc
 accgcccgtc ccgggtatga aactgcacgc gccgtcgtca tcatggcg ggaggagga
 gaaggagaag atgggtata aatgttcggg ttgtgttaag ggattttgggtt ctttatcaagc
 tttaggttgc cacaaagcta gtcacccggaa actcgatccc ggcggagatg atcgtcaac
 tacctccaca accactaacgc caacccggaa aacaacctcc gttAACGGCA acggcaacag
 aagtggaaagg actacgacgt gttcgatttgc tcacaagtgt tttcccactg gacaagctt
 aggtggacac aaaaggtgtc actacgacgg cggtatcggt aacggaaacg ctaacagtgg
 cgtagtgct agcgttggag tgacgtcatc ggaggggtgt gggccacag tcagtccacgg
 ggatttcgac ttgaacattc cggcggttgc ggaattctgg ctgggatttt gttccggcga
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 tgaattatttt caacatataat gggatttga ttgttaggat ttactatttt ggttagacaaa
 attatactat gtaagtttttta attttcatgt tgggtgggag caaaatttt aattttttgt
 ctatagacct agctagttac taatagcaaa aattcaatttgc attgattttaa aaaaaaaaaa
 aaaaaaa

60

120

180

240

300

360

420

480

540

600

660

720

780

840

900

960

1020

1026

<210> 25

<211> 261

<212> PRT

<213> Capsicum annum

<400> 25

Met Ala Leu Glu Ala Leu Asn Ser Pro Thr Gly Thr Pro Thr Pro Pro
 1 5 10 15

Pro Phe Gln Phe Glu Ser Asp Gly Gln Gln Leu Arg Tyr Ile Glu Asn
 20 25 30

Trp Arg Lys Gly Lys Arg Ser Lys Arg Ser Arg Ser Met Glu His Gln

35	40	45	
Pro Thr Glu Glu Glu Tyr Leu Ala Leu Cys Leu Ile Met Leu Ala Arg			
50	55	60	
Ser Gly Gly Ser Val Asn His Gln Arg Ser Leu Pro Pro Pro Ala Pro			
65	70	75	80
Val Met Lys Leu His Ala Pro Ser Ser Ser Ala Ala Glu Glu Glu			
85	90	95	
Lys Glu Lys Met Val Tyr Lys Cys Ser Val Cys Gly Lys Gly Phe Gly			
100	105	110	
Ser Tyr Gln Ala Leu Gly Gly His Lys Ala Ser His Arg Lys Leu Val			
115	120	125	
Pro Gly Gly Asp Asp Gln Ser Thr Thr Ser Thr Thr Thr Asn Ala Thr			
130	135	140	
Gly Thr Thr Thr Ser Val Asn Gly Asn Gly Asn Arg Ser Gly Arg Thr			
145	150	155	160
His Glu Cys Ser Ile Cys His Lys Cys Phe Pro Thr Gly Gln Ala Leu			
165	170	175	
Gly Gly His Lys Arg Cys His Tyr Asp Gly Gly Ile Gly Asn Gly Asn			
180	185	190	
Ala Asn Ser Gly Val Ser Ala Ser Val Gly Val Thr Ser Ser Glu Gly			
195	200	205	
Val Gly Ser Thr Val Ser His Arg Asp Phe Asp Leu Asn Ile Pro Ala			
210	215	220	
Leu Pro Glu Phe Trp Leu Gly Phe Gly Ser Gly Glu Asp Glu Val Glu			
225	230	235	240
Ser Pro His Pro Ala Lys Lys Ser Arg Leu Cys Leu Pro Pro Lys Tyr			
245	250	255	
Glu Leu Phe Gln His			
260			
<210> 26			
<211> 1068			
<212> DNA			
<213> <i>Arabidopsis thaliana</i>			
<400> 26			
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acttttctct caaatttcgt atcctttgtat ccaacagtta gaagaagatt catctgatca			120
tggccctcga aegatgaac actccaactt cttctttcac cagaatcgaa acgaaagaag			180
atttgatgaa cgcgcgggtt ttcattgagc cgtggcttaa acgcaaacgc tccaaacgtc			240
agcgttctca cagcccttct tcgtcttctt cctcaccgcg tcgatctcga cccaaatccc			300
agaatcaaga tcttacggaa gaagagtata tcgcttttg tctcctcatg ctcgctaaag			360
atcaaccgtc gcaaacgcga tttcatcaac agtcgcaatc gttAACGCCG ccGCCAGAAT			420
caaagaacct tccgtacaag tgtaacgtct gtgaaaaagc gtttccttcc tatcaggctt			480

taggcggtca	caaagcaagt	caccgaatca	aaccaccaac	cgtaatctca	acaaccggcg	540
atgattcaac	agctccgacc	atctccatcg	tcgcccggaga	aaaacatccg	attgctgcct	600
ccggaaagat	ccacgagtgt	tcaatctgtc	ataaagtgtt	tccgacgggt	caagcttag	660
gcggtcacaa	acgttgtcac	tacgaaggca	acctcgccgg	cgaggagga	ggaggaagca	720
aatcaatcg	tcacagtgg	agcgtgtcg	gcacggatc	ggaagaaagg	agccaccgtg	780
gattcatcg	tctaaaccta	ccggcggtac	ctgaactcag	ccttcatcac	aatccaatcg	840
tcgacgaaga	gatcttggt	ccgttggacc	gtaaaaaacc	gctttgttg	accgatcact	900
accaagtcat	caagaaaagaa	gatttatctt	taaaaatcta	atactcgact	attaattctt	960
gtgtgatttt	tttcgttaca	accatagttt	cattttcatt	tttttagtta	caaatttta	1020
attgttctga	tttggattga	atattggat	attgttaggg	gttgatac		1068

<210> 27
<211> 273
<212> PRT
<213> *Arabidopsis thaliana*

<400> 27
Met Ala Leu Glu Ala Met Asn Thr Pro Thr Ser Ser Phe Thr Arg Ile
1 5 10 15

Glu Thr Lys Glu Asp Leu Met Asn Asp Ala Val Phe Ile Glu Pro Trp
20 25 30

Leu Lys Arg Lys Arg Ser Lys Arg Gln Arg Ser His Ser Pro Ser Ser
35 40 45

Ser Ser Ser Ser Pro Pro Arg Ser Arg Pro Lys Ser Gln Asn Gln Asp
50 55 60

Leu Thr Glu Glu Glu Tyr Leu Ala Leu Cys Leu Leu Met Leu Ala Lys
65 70 75 80

Asp Gln Pro Ser Gln Thr Arg Phe His Gln Gln Ser Gln Ser Leu Thr
85 90 95

Pro Pro Pro Glu Ser Lys Asn Leu Pro Tyr Lys Cys Asn Val Cys Glu
100 105 110

Lys Ala Phe Pro Ser Tyr Gln Ala Leu Gly Gly His Lys Ala Ser His
115 120 125

Arg Ile Lys Pro Pro Thr Val Ile Ser Thr Thr Ala Asp Asp Ser Thr
130 135 140

Ala Pro Thr Ile Ser Ile Val Ala Gly Glu Lys His Pro Ile Ala Ala
145 150 155 160

Ser Gly Lys Ile His Glu Cys Ser Ile Cys His Lys Val Phe Pro Thr
165 170 175

Gly Gln Ala Leu Gly Gly His Lys Arg Cys His Tyr Glu Gly Asn Leu
180 185 190

Gly Gly Gly Gly Gly Ser Lys Ser Ile Ser His Ser Gly Ser
195 200 205

Val Ser Ser Thr Val Ser Glu Glu Arg Ser His Arg Gly Phe Ile Asp
210 215 220

Leu Asn Leu Pro Ala Leu Pro Glu Leu Ser Leu His His Asn Pro Ile
 225 230 235 240
 Val Asp Glu Glu Ile Leu Ser Pro Leu Thr Gly Lys Lys Pro Leu Leu
 245 250 255
 Leu Thr Asp His Asp Gln Val Ile Lys Lys Glu Asp Leu Ser Leu Lys
 260 265 270

Ile

<210> 28
 <211> 976
 <212> DNA
 <213> *Arabidopsis thaliana*

<400> 28
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 ctgtttcaag attcagcact agggtttcat ggaagcaag gcaaacgatc taagcgatca 180
 agatctgaat tcgaccgtca gagtctcaag gaggatgaat atatcgcttt atgtctcatg 240
 cttcttgcgc gcgacggaga tagaaaccgt gaccttgacc tgcccttcttc ttctgtctca 300
 ctcctctgc ttccctctt tcctactccg atctacaatgt gtagcgtctg tgacaaggcg 360
 ttttcgtctt accaggctct tggtgacac aagcaagtc accggaaaag cttttcgtctt 420
 actcaatctg cggaggaga ttagctgtcg acatcgctgg cgataaccac gtctggata 480
 tccgggtggcg ggggaggaaag tggtaagtcg cacgttgcg ctatctgtca taaatcggtc 540
 gcccacggtc aagctctcgcc cggccacaaa cggtgccact acgaaggaaa gaacggaggc 600
 ggtgtagta gtagcgtgtc gaattctgaa gatgtggggct ctacaagccca cgtcagcagt 660
 ggccacccgtg ggttgacact caacataccg cggataccgg aattctcgat ggtcaacgg 720
 gacgaagagg tggatggatcc tatggccggcg aagaaactcc ggttgactt cccggagaaa 780
 ccctaaacat aaacacttaga aaaactttac agaatttcatt ttataggaaa ttgttttact 840
 gtatatacaa atatcgattt tgattgtatg tcttcac tggaaaattt tgattcttg 900
 ttgtataatt gatgttctg aaaaagatata aacttttat tgttcacac gtatcaaaat 960
 ttgttgcgtt acatca 976

<210> 29
 <211> 238
 <212> PRT
 <213> *Arabidopsis thaliana*

<400> 29
 Met Ala Leu Glu Thr Leu Thr Ser Pro Arg Leu Ser Ser Pro Met Pro
 1 5 10 15
 Thr Leu Phe Gln Asp Ser Ala Leu Gly Phe His Gly Ser Lys Gly Lys
 20 25 30
 Arg Ser Lys Arg Ser Arg Ser Glu Phe Asp Arg Gln Ser Leu Thr Glu
 35 40 45
 Asp Glu Tyr Ile Ala Leu Cys Leu Met Leu Leu Ala Arg Asp Gly Asp
 50 55 60
 Arg Asn Arg Asp Leu Asp Leu Pro Ser Ser Ser Ser Pro Pro Leu
 65 70 75 80

Leu Pro Pro Leu Pro Thr Pro Ile Tyr Lys Cys Ser Val Cys Asp Lys
85 90 95

Ala Phe Ser Ser Tyr Gln Ala Leu Gly Gly His Lys Ala Ser His Arg
100 105 110

Lys Ser Phe Ser Leu Thr Gln Ser Ala Gly Gly Asp Glu Leu Ser Thr
115 120 125

Ser Ser Ala Ile Thr Thr Ser Gly Ile Ser Gly Gly Gly Ser
130 135 140

Val Lys Ser His Val Cys Ser Ile Cys His Lys Ser Phe Ala Thr Gly
145 150 155 160

Gln Ala Leu Gly Gly His Lys Arg Cys His Tyr Glu Gly Lys Asn Gly
165 170 175

Gly Gly Val Ser Ser Ser Val Ser Asn Ser Glu Asp Val Gly Ser Thr
180 185 190

Ser His Val Ser Ser Gly His Arg Gly Phe Asp Leu Asn Ile Pro Pro
195 200 205

Ile Pro Glu Phe Ser Met Val Asn Gly Asp Glu Glu Val Met Ser Pro
210 215 220

Met Pro Ala Lys Lys Leu Arg Phe Asp Phe Pro Glu Lys Pro
225 230 235

<210> 30

<211> 718

<212> DNA

<213> *Arabidopsis thaliana*

<400> 30

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ttcctccgtt gcctcgacga aaccgagccc gaaaacctcg aatcatggac caaaagaaaa
cgtacaaaac gtcaccgtat agatcaacca aaccctcctc cttctgaaaga agagtatctc
gctcttgc tccttatgct cgctcggtgc tcctccgatc atcactctcc accgtcgat
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aaatcttcc cgtcttacca agcgttagt ggacacaaaa caagtcaccg gaaaccgggt
agtgtcgatg ttaataatag taacggaaacc gttactaata acggaaatat tagtaacgg
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ggtcaagcat tgggtggta caaacgttgt cactatgatg gtggtaacgg taacagtaac
ggtacaataa gcccacaagtt tgacctaattt acggctgt atcaagttatg tgatgagaca
attggaaaaaa gtcaactctc cggtaagaa acaaagtcgg tgggtgtattt attattattt
tttacggatc gggatttagt agtgggtat cattagctga gtctgtaatg aaaatgtat

60

120

180

240

300

360

420

480

540

600

660

718

<210> 31

<211> 215

<212> PRT

<213> *Arabidopsis thaliana*

<400> 31

Met Ala Leu Asp Thr Leu Asn Ser Pro Thr Ser Thr Thr Thr Thr
1 5 10 15

Ala Pro Pro Pro Phe Leu Arg Cys Leu Asp Glu Thr Glu Pro Glu Asn
 20 25 30

Leu Glu Ser Trp Thr Lys Arg Lys Arg Thr Lys Arg His Arg Ile Asp
 35 40 45

Gln Pro Asn Pro Pro Ser Glu Glu Glu Tyr Leu Ala Leu Cys Leu
 50 55 60

Leu Met Leu Ala Arg Gly Ser Ser Asp His His Ser Pro Pro Ser Asp
 65 70 75 80

His His Ser Leu Ser Pro Leu Ser Asp His Gln Lys Asp Tyr Lys Cys
 85 90 95

Ser Val Cys Gly Lys Ser Phe Pro Ser Tyr Gln Ala Leu Gly Gly His
 100 105 110

Lys Thr Ser His Arg Lys Pro Val Ser Val Asp Val Asn Asn Ser Asn
 115 120 125

Gly Thr Val Thr Asn Asn Gly Asn Ile Ser Asn Gly Leu Val Gly Gln
 130 135 140

Ser Gly Lys Thr His Asn Cys Ser Ile Cys Phe Lys Ser Phe Pro Ser
 145 150 155 160

Gly Gln Ala Leu Gly Gly His Lys Arg Cys His Tyr Asp Gly Gly Asn
 165 170 175

Gly Asn Ser Asn Gly Asp Asn Ser His Lys Phe Asp Leu Asn Leu Pro
 180 185 190

Ala Asp Gln Val Ser Asp Glu Thr Ile Gly Lys Ser Gln Leu Ser Gly
 195 200 205

Glu Glu Thr Lys Ser Val Leu
 210 215

<210> 32
 <211> 702
 <212> DNA
 <213> *Arabidopsis thaliana*

<400> 32
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 ttaagattca atggcggttga gcagtggacc aaatgttaga aacgtatccaa acgttcgaga 120
 tctgtatcttc atcataacca ccgttctcaact gaggaagagt atcttagctt ctgtctcatg 180
 cttcttgctc gggatggcg cgatcttgcac tctgtgacgg ttgcggagaa gcccggat 240
 aagtgtggcg tttgttacaa gacgttttcg tcttaccaag ctctcgccgg tcataaagcg 300
 agccaccgga gcttatacgg tggtggagag aatgataaaat cgacaccatc caccggccgtg 360
 aaatctcacg tttgttcggt ttgcgggaaa tctttcgcca ccgggtcaagc tctcgccggc 420
 cacaaggcggt gccactacga tggtggcggt tcgaactcgg aagggtgtggg gtctactagc 480
 cacgtcagca gtatgttacca ccgtggattt gacctaata ttataccggt gcagggattt 540
 tcgcccggacg acgaagtgtat gagtccgtt gcgactaaga agcctcgccct gaagtaagtc 600
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 ttttggat tctattacca atacacaata cgattcaattt cc 702

<210> 33
 <211> 193
 <212> PRT
 <213> *Arabidopsis thaliana*

<400> 33
 Met Ala Leu Glu Ala Leu Asn Ser Pro Arg Leu Val Glu Asp Pro Leu
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 Arg Phe Asn Gly Val Glu Gln Trp Thr Lys Cys Lys Lys Arg Ser Lys
 20 25 30
 Arg Ser Arg Ser Asp Leu His His Asn His Arg Leu Thr Glu Glu
 35 40 45
 Tyr Leu Ala Phe Cys Leu Met Leu Leu Ala Arg Asp Gly Gly Asp Leu
 50 55 60
 Asp Ser Val Thr Val Ala Glu Lys Pro Ser Tyr Lys Cys Gly Val Cys
 65 70 75 80
 Tyr Lys Thr Phe Ser Ser Tyr Gln Ala Leu Gly Gly His Lys Ala Ser
 85 90 95
 His Arg Ser Leu Tyr Gly Gly Glu Asn Asp Lys Ser Thr Pro Ser
 100 105 110
 Thr Ala Val Lys Ser His Val Cys Ser Val Cys Gly Lys Ser Phe Ala
 115 120 125
 Thr Gly Gln Ala Leu Gly Gly His Lys Arg Cys His Tyr Asp Gly Gly
 130 135 140
 Val Ser Asn Ser Glu Gly Val Gly Ser Thr Ser His Val Ser Ser Ser
 145 150 155 160
 Ser His Arg Gly Phe Asp Leu Asn Ile Ile Pro Val Gln Gly Phe Ser
 165 170 175
 Pro Asp Asp Glu Val Met Ser Pro Met Ala Thr Lys Lys Pro Arg Leu
 180 185 190
 Lys

<210> 34
 <211> 1157
 <212> DNA
 <213> *Arabidopsis thaliana*

<400> 34
 cacacttcac tctttcttca tctttcttctt cttaaatagc tcgaaatcac atctcacaga 60
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 tcttctccgg tategtgaag aaatggagcc tgagaatctc gagcaatggg ctaaaagaaa 180
 acgaacaaaa cgtcaacgtt ttgatcacgg tcatacagaat caagaaacga acaagaacct 240
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 acaatctctt cctcttcctc ctctaccgtc acgtgcgtca ccgtccgatc accgagatta 360
 caagtgtacg gtctgtggga agtccttttc gtcataccaa gccttaggtg gacacaagac 420

gagtcacccgg	aaaccgacga	acaactagtat	caacttccgg	aaccaagaac	tgtctaataa	480
cagtcacagt	aacagcggtt	ccgttgttat	taacgttacc	gtgaacactg	gtaacggtgt	540
tagtcaaagc	ggaaagattc	acaacttgctc	aatctgttcc	aagtcggttg	cgtctggtca	600
agccttaggt	ggacacaaac	ggtgtcacta	tgacgggtggc	aacaacggta	acggtAACGG	660
aagtagcagc	aacagcgtag	aactcgtcgc	tggttagtgac	gtcagcgtat	ttgataatga	720
gagatggccc	gaagaaaatgt	cgatcggtgg	ccaccgtgga	tttgacctaactt	tttaccggc	780
tgatcaagtc	tcagtacga	cttcttaac	ttgactgagt	ttgaggaaaa	agtcaactat	840
caagcgaaga	aagggttagt	ggacgggtgaa	gattaacgg	cgtttcttc	cagttgcttc	900
ggttttagct	tgactgggtc	tgtaatgaaa	atgattggag	tggacttggc	attattatttt	960
ttatattttaa	aaagaaatgt	taatttgttg	ttggatttgt	ttatagatag	aggaaaacaat	1020
tgggatacac	aaatattttt	tttttttaca	aagaaaataa	taatgcagag	atggatgatt	1080
ggatcgtaca	cgtttattata	tagtggacca	ttctgtatc	gtgaattattt	attattttgtt	1140
agaaaattttaa	ttttcg					1157

<210> 35

<211> 245

<212> PRT

<213> *Arabidopsis thaliana*

<400> 35

Met	Ala	Leu	Glu	Thr	Leu	Asn	Ser	Pro	Thr	Ala	Thr	Thr	Ala	Arg
1					5				10				15	

Pro	Leu	Leu	Arg	Tyr	Arg	Glu	Glu	Met	Glu	Pro	Glu	Asn	Leu	Glu	Gln
								20	25				30		

Trp	Ala	Lys	Arg	Lys	Arg	Thr	Lys	Arg	Gln	Arg	Phe	Asp	His	Gly	His
							35	40				45			

Gln	Asn	Gln	Glu	Thr	Asn	Lys	Asn	Leu	Pro	Ser	Glu	Glu	Glu	Tyr	Leu
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Ala	Leu	Cys	Leu	Leu	Met	Leu	Ala	Arg	Gly	Ser	Ala	Val	Gln	Ser	Pro
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Pro	Leu	Pro	Pro	Leu	Pro	Ser	Arg	Ala	Ser	Pro	Ser	Asp	His	Arg	Asp
						85	90				95				

Tyr	Lys	Cys	Thr	Val	Cys	Gly	Lys	Ser	Phe	Ser	Ser	Tyr	Gln	Ala	Leu
						100		105				110			

Gly	Gly	His	Lys	Thr	Ser	His	Arg	Lys	Pro	Thr	Asn	Thr	Ser	Ile	Thr
						115		120			125				

Ser	Gly	Asn	Gln	Glu	Leu	Ser	Asn	Asn	Ser	His	Ser	Asn	Ser	Gly	Ser
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Val	Val	Ile	Asn	Val	Thr	Val	Asn	Thr	Gly	Asn	Gly	Val	Ser	Gln	Ser
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Gly	Lys	Ile	His	Thr	Cys	Ser	Ile	Cys	Phe	Lys	Ser	Phe	Ala	Ser	Gly
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Gln	Ala	Leu	Gly	Gly	His	Lys	Arg	Cys	His	Tyr	Asp	Gly	Gly	Asn	Asn
						180		185			190				

Gly	Asn	Gly	Asn	Gly	Ser	Ser	Ser	Asn	Ser	Val	Glu	Leu	Val	Ala	Gly
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Ser Asp Val Ser Asp Val Asp Asn Glu Arg Trp Ser Glu Glu Ser Ala
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Ile Gly Gly His Arg Gly Phe Asp Leu Asn Leu Pro Ala Asp Gln Val
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Ser Val Thr Thr Ser
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<212> DNA

<213> Oryza sativa

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<212> PRT

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Ser Ser Ala Thr Ser Gly Glu Glu Gly Gly His Leu Pro Gln Gly Trp		
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Ala Lys Arg Lys Arg Ser Arg Arg Gln Arg Ser Glu Glu Glu Asn Leu		
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Ala Leu Cys Leu Leu Met Leu Ala Arg Gly Gly His His Arg Val Gln		
65	70	80

Ala Pro Pro Pro Leu Ser Ala Ser Ala Pro Pro Pro Ala Gly Ala Glu
 85 90 95

Phe Lys Cys Ser Val Cys Gly Lys Ser Phe Ser Ser Tyr Gln Ala Leu
 100 105 110

Gly Gly His Lys Thr Ser His Arg Val Lys Leu Pro Thr Pro Pro Ala
 115 120 125

Ala Pro Val 'Leu Ala Pro Ala Pro Val Ala Ala Leu Leu Pro Ser Ala
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Glu Asp Arg Glu Pro Ala Thr Ser Ser Thr Ala Ala Ser Ser Asp Gly
 145 150 155 160

Met Thr Asn Arg Val His Arg Cys Ser Ile Cys Gln Lys Glu Phe Pro
 165 170 175

Thr Gly Gln Ala Leu Gly Gly His Lys Arg Lys His Tyr Asp Gly Gly
 180 185 190

Val Gly Ala Gly Ala Gly Ala Ser Ser Thr Glu Leu Leu Ala Thr Val
 195 200 205

Ala Ala Glu Ser Glu Val Gly Ser Ser Gly Asn Gly Gln Ser Ala Thr
 210 215 220

Arg Ala Phe Asp Leu Asn Leu Pro Ala Val Pro Glu Phe Val Trp Arg
 225 230 235 240

Pro Cys Ser Lys Gly Lys Lys Met Trp Asp Glu Glu Glu Val Gln
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Ser Pro Leu Ala Phe Lys Lys Pro Arg Leu Leu Thr Ala
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<210> 38

<211> 528

<212> DNA

<213> *Arabidopsis thaliana*

<400> 38

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Ile Asp Val Lys Gln Ser Thr Gly Ser Lys Thr Asn His Asn Asn His
35 40 45

Phe Glu Cys Lys Thr Cys Asn Arg Lys Phe Asp Ser Phe Gln Ala Leu
50 55 60

Gly Gly His Arg Ala Ser His Lys Lys Pro Lys Leu Ile Val Asp Gln
65 70 75 80

Glu Gln Val Lys His Arg Asn Lys Glu Asn Asp Met His Lys Cys Thr
85 90 95

Ile Cys Asp Gln Met Phe Gly Thr Gly Gln Ala Leu Gly Gly His Met
100 105 110

Arg Lys His Arg Thr Ser Met Ile Thr Glu Gln Ser Ile Val Pro Ser
115 120 125

Val Val Tyr Ser Arg Pro Val Phe Asn Arg Cys Ser Ser Ser Lys Glu
130 135 140

Ile Leu Asp Leu Asn Leu Thr Pro Leu Glu Asn Asp Leu Val Leu Ile
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Phe Gly Lys Asn Leu Val Pro Gln Ile Asp Leu Lys Phe Val Asn
165 170 175

<210> 40

<211> 820

<212> DNA

<213> *Saccharum officinarum*

<220>

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<222> (406)..(406)

<223> n can be any nucleotide

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<223> n can be any nucleotide

<400> 40

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<211> 1509

<212> DNA

<213> *Arabidopsis thaliana*

<400> 41

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<210> 42

<211> 383

<212> PRT

<213> *Arabidopsis thaliana*

<400> 42

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20							25							30		

Leu Pro Pro Val Thr Pro Pro Ser Ser Phe Phe Phe Phe Pro Gln Ser
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Gly Asp Leu Arg Arg Pro Pro Pro Pro Pro Pro Thr Pro Pro Pro Ser Pro
50 55 60

Pro Leu Arg Glu Ala Leu Pro Leu Leu Ser Leu Ser Pro Ala Asn Lys
65 70 75 80

Gln Gln Asp His His His Asn His Asp His Leu Ile Gln Glu Pro Pro
85 90 95

Ser Thr Ser Met Asp Val Asp Tyr Asp His His His Gln Asp Asp His
100 105 110

His Asn Leu Asp Asp Asp His Asp Val Thr Val Ala Leu His Ile
115 120 125

Gly Leu Pro Ser Pro Ser Ala Gln Glu Met Ala Ser Leu Leu Met Met
130 135 140

Ser Ser Ser Ser Ser Ser Arg Thr Thr His His His Glu Asp Met
145 150 155 160

Asn His Lys Lys Asp Leu Asp His Glu Tyr Ser His Gly Ala Val Gly
165 170 175

Gly Gly Glu Asp Asp Asp Glu Asp Ser Val Gly Gly Asp Gly Gly Cys
180 185 190

Arg Ile Ser Arg Leu Asn Lys Gly Gln Tyr Trp Ile Pro Thr Pro Ser
195 200 205

Gln Ile Leu Ile Gly Pro Thr Gln Phe Ser Cys Pro Val Cys Phe Lys
210 215 220

Thr Phe Asn Arg Tyr Asn Asn Met Gln Met His Met Trp Gly His Gly
225 230 235 240

Ser Gln Tyr Arg Lys Gly Pro Glu Ser Leu Arg Gly Thr Gln Pro Thr
245 250 255

Gly Met Leu Arg Leu Pro Cys Tyr Cys Cys Ala Pro Gly Cys Arg Asn
260 265 270

Asn Ile Asp His Pro Arg Ala Lys Pro Leu Lys Asp Phe Arg Thr Leu
275 280 285

Gln Thr His Tyr Lys Arg Lys His Gly Ile Lys Pro Phe Met Cys Arg
290 295 300

Lys Cys Gly Lys Ala Phe Ala Val Arg Gly Asp Trp Arg Thr His Glu
305 310 315 320

Lys Asn Cys Gly Lys Leu Trp Tyr Cys Ile Cys Gly Ser Asp Phe Lys
325 330 335

His Lys Arg Ser Leu Lys Asp His Ile Lys Ala Phe Gly Asn Gly His

340 345 350

Gly Ala Tyr Gly Ile Asp Gly Phe Asp Glu Glu Asp Glu Pro Ala Ser
 355 360 365

Glu Val Glu Gln Leu Asp Asn Asp His Glu Ser Met Gln Ser Lys
 370 375 380

<210> 43

<211> 1303

<212> DNA

<213> *Arabidopsis thaliana*

<400> 43

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 atacgggtct ttatattcttgc atcttcttctc tcttctttgt ctctccctca gagtcatgtc
 taatccagct tgttcgaatc tcttcaacaa tggatgtgac cataataatgc tcaactattc
 cacttcttc tcttacattt acaactctca cggtagtac tattactcta ataccacaaa
 ccctaattac attaatcata ctcataaccac ttccacttcc cctaactcac ccccaacta
 agaagcttt cctcttctta gcttaagccc cataaggcac caagaacaac aagaccaaca
 ctatccatg gacacccatc aaattagctc ttcaaaactt ctgtatgtac ctcttgc
 tgtagatctt catctagggt taccaaacta cggtaggtt gtagacattt ggagcaatat
 tgctcctgtat gcaaccacgg acggacaa tcaagatcat gaccgaggag tagaagtac
 agttgatcc cacccatc atgatgtatc tcatcatgaa gatctacaca gaggtcatca
 ctatccatg cctactcctt ctcagatttt gattggctt acacagtca ctgttgc
 ttgcttcaag acattcaaca gatacaacaa catgcagatg cacatgtgg gacacggctc
 acaatacaga aaggacccatc aatccttaag aggaacccaa ccaacaggaa tgctaaact
 accatgtttt tgctgtgcac ccgggtgcaaa gaacaacatt gaccacccac gagccaagcc
 tcttaaggac tttcaacacc tccaaacaca ttacaaacgt aaacatgggt ctaaaccatt
 tgcttgcgt atgtgtggta aggcctttgc agtgaaggaa gattggagaa cgcatgagaa
 gaatttgaa aagctttggat attgctctt tggctcgat tttaaagcaca agaggtcgct
 taaggacat gtcaaggcct ttggaaatgg tcatgttct tggggattt atagtttgg
 agggatcat gaggactact atgatgtgc ttctgatatc gagaataag atgatagcaa
 caacaatgag tgtaatttag gggttttgtt tattttctt ctcatgcatt agttgattgt
 atgcacgtgt tcttttagttt tgttcttcgg atctttgtt tattttgtt tgagctgttt
 ttttttaat tactaagaag ttaattatca tctaaagatt ttc 1303

<210> 44

<211> 337

<212> PRT

<213> *Arabidopsis thaliana*

<400> 44

Met Ser Asn Pro Ala Cys Ser Asn Leu Phe Asn Asn Gly Cys Asp His
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Asn Ser Phe Asn Tyr Ser Thr Ser Leu Ser Tyr Ile Tyr Asn Ser His
 20 25 30

Gly Ser Tyr Tyr Tyr Ser Asn Thr Thr Asn Pro Asn Tyr Ile Asn His
 35 40 45

Thr His Thr Thr Ser Thr Ser Pro Asn Ser Pro Pro Leu Arg Glu Ala
 50 55 60

Leu Pro Leu Leu Ser Leu Ser Pro Ile Arg His Gln Glu Gln Gln Asp
 65 70 75 80

Gln His Tyr Phe Met Asp Thr His Gln Ile Ser Ser Ser Asn Phe Leu
 85 90 95
 Asp Asp Pro Leu Val Thr Val Asp Leu His Leu Gly Leu Pro Asn Tyr
 100 105 110
 Gly Val Gly Glu Ser Ile Arg Ser Asn Ile Ala Pro Asp Ala Thr Thr
 115 120 125
 Asp Glu Gln Asp Gln Asp His Asp Arg Gly Val Glu Val Thr Val Glu
 130 135 140
 Ser His Leu Asp Asp Asp Asp His His Gly Asp Leu His Arg Gly
 145 150 155 160
 His His Tyr Trp Ile Pro Thr Pro Ser Gln Ile Leu Ile Gly Pro Thr
 165 170 175
 Gln Phe Thr Cys Pro Leu Cys Phe Lys Thr Phe Asn Arg Tyr Asn Asn
 180 185 190
 Met Gln Met His Met Trp Gly His Gly Ser Gln Tyr Arg Lys Gly Pro
 195 200 205
 Glu Ser Leu Arg Gly Thr Gln Pro Thr Gly Met Leu Arg Leu Pro Cys
 210 215 220
 Phe Cys Cys Ala Pro Gly Cys Lys Asn Asn Ile Asp His Pro Arg Ala
 225 230 235 240
 Lys Pro Leu Lys Asp Phe Arg Thr Leu Gln Thr His Tyr Lys Arg Lys
 245 250 255
 His Gly Ser Lys Pro Phe Ala Cys Arg Met Cys Gly Lys Ala Phe Ala
 260 265 270
 Val Lys Gly Asp Trp Arg Thr His Glu Lys Asn Cys Gly Lys Leu Trp
 275 280 285
 Tyr Cys Ser Cys Gly Ser Asp Phe Lys His Lys Arg Ser Leu Lys Asp
 290 295 300
 His Val Lys Ala Phe Gly Asn Gly His Val Pro Cys Gly Ile Asp Ser
 305 310 315 320
 Phe Gly Gly Asp His Glu Asp Tyr Tyr Asp Ala Ala Ser Asp Ile Glu
 325 330 335

Gln

<210> 45
 <211> 495
 <212> DNA
 <213> *Arabidopsis thaliana*

<400> 45
 atggttgcga gaagtgagga agttgagata gtggaagata cggcggcgaa atgtttgatg

ttgttatcaa gagttggaga atgcggcgg a gaggagaga aacgagttt ccgatgcaag 120
 acttgtctt a aaggtttc gtcgttcaa gctttggag gtcatcgac aagccacaag 180
 aaactcatta acatgtcgatccatctt cttggatctt tgcataacaa gaaaactaaa 240
 acggcgacgt ctcatcccttgcgtatgt ggcgtggat ttccgatggg gcaagcttt 300
 ggtggtcaca tgaggagaca taggagttagaa gaagcctcac caggcacgtt ggttacacgt 360
 tcttttttac cggagacgac gacggtgacg acttggaaaa aatcgagtag tggaaagaga 420
 gtggcttggg tggacttaga ttgcgtggag agtttagtca attggaaagtt ggagttggga 480
 agaacgattt ctgta 495

<210> 46
 <211> 164
 <212> PRT
 <213> *Arabidopsis thaliana*

<400> 46
 Met Val Ala Arg Ser Glu Glu Val Glu Ile Val Glu Asp Thr Ala Ala
 1 5 10 15

Lys Cys Leu Met Leu Leu Ser Arg Val Gly Glu Cys Gly Gly Gly
 20 25 30

Glu Lys Arg Val Phe Arg Cys Lys Thr Cys Leu Lys Glu Phe Ser Ser
 35 40 45

Phe Gln Ala Leu Gly Gly His Arg Ala Ser His Lys Lys Leu Ile Asn
 50 55 60

Ser Ser Asp Pro Ser Leu Leu Gly Ser Leu Ser Asn Lys Lys Thr Lys
 65 70 75 80

Thr Ala Thr Ser His Pro Cys Pro Ile Cys Gly Val Glu Phe Pro Met
 85 90 95

Gly Gln Ala Leu Gly Gly His Met Arg Arg His Arg Ser Glu Lys Ala
 100 105 110

Ser Pro Gly Thr Leu Val Thr Arg Ser Phe Leu Pro Glu Thr Thr Thr
 115 120 125

Val Thr Thr Leu Lys Lys Ser Ser Ser Gly Lys Arg Val Ala Cys Leu
 130 135 140

Asp Leu Asp Ser Met Glu Ser Leu Val Asn Trp Lys Leu Glu Leu Gly
 145 150 155 160

Arg Thr Ile Ser

<210> 47
 <211> 1209
 <212> DNA
 <213> *Arabidopsis thaliana*

<400> 47
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 attactcggt cacagaaaga aacagaaaag tctacgaacc aacagcaaga tggacttgt 120
 tactatggtc taaggaaaa ctcgaagaag aaaacccagg aatctccggg accaatgaag 180
 aagattttgtt ttcgtgcga agaatgtgaa aaagggttgc ggtacgagaa atattttaa 240

aatcatcgct	cgatgatgca	tttatcgccg	aacgagaagg	tttgtgaaga	atccttgatg	300
actctgtctc	gtagccttgg	gtttgtgaag	aagaagaaaa	gatcaagact	tggtaggtct	360
gggaagactt	tatttactac	gtttcttga	ccgagttcta	tttttgcata	gactgatgaa	420
gaattagaag	tggcggttgg	tttgattcta	ttgtctaaaga	gtgctcccaa	ggtttagac	480
gaattgaaaa	gtcttctga	ggcagttacgt	gttactcctg	aaacaccctga	aagttagctat	540
gatttgggtt	gtttgtctaa	caagaaaccg	agaaaagggtg	gtgaaatttgg	atctgggggtt	600
ttaagtaatg	agcaaagact	tatggaaagaa	gggttttagta	gttatggaaac	atcgaaaagaa	660
ccagctagct	tcttgagaga	cggaaacaga	ttggatcagc	agaaacggag	aaaagatgg	720
gaatttgaat	ccggactttt	gatgtatgg	caaagactgc	tagaagaaga	gattactact	780
cctgtgacat	tcaaaaggccc	agcgagttcc	ttgagacaca	agtgtgcctt	ggatcgaaat	840
ggaggtgaat	ttggctctga	gtttttgagt	aatgagcaaa	cactgtatgg	agaaacatgg	900
aaagaaccag	tgagtttctt	agaagataag	catgaatttg	atcagcggaa	aatgcgagaa	960
gctggcgact	ttgaatctag	gttttacaga	attgagcttg	gagtaggagc	tatggagttgt	1020
acttcttcag	atactgatata	gctcacgca	tctgataaga	agaacgttga	gcatcgatgc	1080
aggttgtgca	acaagatatt	ctcgctttat	caagctctag	gggttcatca	gacgtttcat	1140
cgatgagca	aatgtaaagaa	caagaagaat	ggcatagagg	aatcagtta	acccaggatg	1200
actctgtga						1209

<210> 48

<211> 402

<212> PRT

<213> *Arabidopsis thaliana*

<400> 48

Met	Glu	Asp	Glu	His	Gln	Asp	Leu	His	Lys	Pro	Ile	Asn	Gly	Ala	Leu
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Arg	Asp	Leu	Lys	Ile	Thr	Arg	Ser	Gln	Lys	Glu	Thr	Glu	Lys	Ser	Thr
															20
															25

Asn	Gln	Gln	Gln	Asp	Val	Thr	Cys	Tyr	Tyr	Gly	Leu	Arg	Glu	Asn	Ser
															35
															40

Lys	Lys	Lys	Thr	Gln	Glu	Ser	Pro	Glu	Pro	Met	Lys	Lys	Ile	Leu	Phe
															50
															55

Arg	Cys	Glu	Glu	Cys	Gly	Lys	Gly	Phe	Arg	Tyr	Glu	Lys	Tyr	Phe	Lys
															65
															70

Asn	His	Arg	Ser	Met	Met	His	Leu	Ser	Pro	Asn	Glu	Lys	Val	Cys	Glu
															85
															90

Glu	Ser	Leu	Met	Thr	Leu	Ser	Arg	Ser	Leu	Gly	Phe	Val	Lys	Lys	Lys
															100
															105

Lys	Arg	Ser	Arg	Leu	Gly	Arg	Ser	Gly	Lys	Thr	Leu	Phe	Thr	Thr	Phe
															115
															120

Leu	Glu	Pro	Ser	Ser	Ile	Phe	Asp	Ala	Thr	Asp	Glu	Glu	Leu	Glu	Val
															130
															135

Ala	Asp	Cys	Leu	Ile	Leu	Ser	Lys	Ser	Ala	Pro	Lys	Val	Val	Asp	
															145
															150

Glu	Leu	Lys	Ser	Leu	Ser	Glu	Ala	Val	Arg	Val	Thr	Pro	Glu	Thr	Pro
															165
															170

Glu	Ser	Ser	Tyr	Asp	Leu	Gly	Cys	Leu	Leu	Asn	Lys	Lys	Pro	Arg	Lys
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180	185	190
Gly Gly Glu Leu Glu Ser Gly Val Leu Ser Asn Glu Gln Arg Leu Met		
195	200	205
Glu Glu Gly Phe Ser Ser Tyr Gly Thr Ser Lys Glu Pro Ala Ser Phe		
210	215	220
Leu Arg Asp Glu Asn Arg Leu Asp Gln Gln Lys Arg Arg Lys Asp Gly		
225	230	235
Glu Phe Glu Ser Gly Leu Leu Ser Asn Glu Gln Arg Leu Leu Glu Glu		
245	250	255
Glu Ile Thr Thr Pro Val Thr Phe Lys Gly Pro Ala Ser Ser Leu Arg		
260	265	270
His Lys Cys Ala Leu Asp Arg Asn Gly Gly Glu Phe Gly Pro Glu Phe		
275	280	285
Leu Ser Asn Glu Gln Thr Leu Met Glu Glu Thr Trp Lys Glu Pro Val		
290	295	300
Ser Phe Leu Glu Asp Lys His Glu Phe Asp Gln Arg Lys Met Arg Glu		
305	310	320
Ala Gly Asp Phe Glu Ser Arg Phe Tyr Arg Ile Glu Leu Gly Val Gly		
325	330	335
Ala Met Glu Cys Thr Ser Ser Asp Thr Asp Met Leu Thr Gln Ser Asp		
340	345	350
Lys Lys Asn Val Glu His Arg Cys Arg Leu Cys Asn Lys Ile Phe Ser		
355	360	365
Ser Tyr Gln Ala Leu Gly Gly His Gln Thr Phe His Arg Met Ser Lys		
370	375	380
Cys Lys Asn Lys Lys Asn Gly Ile Glu Glu Ser Val Glu Pro Arg Met		
385	390	395
Thr Leu		

<210> 49
 <211> 1087
 <212> DNA
 <213> *Arabidopsis thaliana*

<400> 49
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 aagtttcttg aattgggttt gtttcttgc ttgtttcttg aattgggttt tggctttctt 120
 ttcttactat atttggatat gatgatgggt caagatgggg ttgggagtga tcagacgcaa 180
 atcataaaag ggaaacgtac gaagcgacaa agatcgctt cgacgtttgt ggtgacggcg 240
 gcgacaacag tgacttcaac aagttcatcg gccggtgaa gtggaggaga aagagctgtt 300
 tcagatgaaat acaactcgcc ggtttcgctt ccgggtacta ctgattgtac gcaagaagaa 360
 gaagacatgg cgatttgcgtt catcatgtta gctcggtggaa cagttcttcc atcgccggat 420
 ctcaagaact cgagaaaaat tcatcagaag atttcgctgg agaattcttag tttctatgtg 480

tacgagtgtta	aaacgtgtaa	cggacgttt	tcgtcggtcc	aagcacttgg	tggacacaga	540
gcgagccaca	agaagccgag	gacgtcgact	gaggaaaaga	ctagactacc	cctgacgcaa	600
cccaagtcta	gtgcatcaga	agaagggcaa	aacagtctt	tcaaagtttc	cggctcagcc	660
ctagcttac	aggcaagtaa	catcatcaac	aaggcaaaca	aagtacacga	gtgttccatc	720
tgcggttctg	agttcacttc	cgggcaagct	ctcggtggtc	acatgaggcg	gcacaggaca	780
gccccaaacca	cgattagccc	cgttgcagcc	accgcagaag	taagcagaaa	cagtacagag	840
gaagagatg	agatcaat	aggccgttctg	atggacagc	agagggaaata	tctaccgttg	900
gatcttaatc	taccgcacc	aggagatgt	ctaagagatg	ccaagttca	aggatagta	960
ttctcagcaa	caccagcgtt	aatagattgt	cattactagt	tgtttttttt	actacataat	1020
atgatgaaat	atttgtaat	tcttcttact	tactactata	ttgttgatca	aaaaaaaaaa	1080
aaaaaaaaa						1087

<210> 50

<211> 284

<212> PRT

<213> *Arabidopsis thaliana*

<400> 50

Met Gly Gln Asp Glu Val Gly Ser Asp Gln Thr Gln Ile Ile Lys Gly

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Lys Arg Thr Lys Arg Gln Arg Ser Ser Ser Thr Phe Val Val Thr Ala

20 25 30

Ala Thr Thr Val Thr Ser Thr Ser Ser Ala Gly Gly Ser Gly Gly

35 40 45

Glu Arg Ala Val Ser Asp Glu Tyr Asn Ser Ala Val Ser Ser Pro Val

50 55 60

Thr Thr Asp Cys Thr Gln Glu Glu Asp Met Ala Ile Cys Leu Ile

65 70 75 80

Met Leu Ala Arg Gly Thr Val Leu Pro Ser Pro Asp Leu Lys Asn Ser

85 90 95

Arg Lys Ile His Gln Lys Ile Ser Ser Glu Asn Ser Ser Phe Tyr Val

100 105 110

Tyr Glu Cys Lys Thr Cys Asn Arg Thr Phe Ser Ser Phe Gln Ala Leu

115 120 125

Gly Gly His Arg Ala Ser His Lys Lys Pro Arg Thr Ser Thr Glu Glu

130 135 140

Lys Thr Arg Leu Pro Leu Thr Gln Pro Lys Ser Ser Ala Ser Glu Glu

145 150 155 160

Gly Gln Asn Ser His Phe Lys Val Ser Gly Ser Ala Leu Ala Ser Gln

165 170 175

Ala Ser Asn Ile Ile Asn Lys Ala Asn Lys Val His Glu Cys Ser Ile

180 185 190

Cys Gly Ser Glu Phe Thr Ser Gly Gln Ala Leu Gly Gly His Met Arg

195 200 205

Arg His Arg Thr Ala Val Thr Ile Ser Pro Val Ala Ala Thr Ala

210 215 220

Glu Val Ser Arg Asn Ser Thr Glu Glu Glu Ile Glu Ile Asn Ile Gly
225 230 235 240

Arg Ser Met Glu Gln Gln Arg Lys Tyr Leu Pro Leu Asp Leu Asn Leu
245 250 255

Pro Ala Pro Gly Asp Asp Leu Arg Glu Ser Lys Phe Gln Gly Ile Val
260 265 270

Phe Ser Ala Thr Pro Ala Leu Ile Asp Cys His Tyr
275 280